PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:

C07K 13/00, C12N 15/31, 15/62;
A61K 37/02, C07K 3/18

(11) International Publication Number: WO 93/22342

(43) International Publication Date: 11 November 1993 (11.11.93)

SE

(21) International Application Number: PCT/SE93/00375

(22) International Filing Date: 28 April 1993 (28.04.93)

(22) International Fining Date: 28 April 1993 (28.04.93)

28 April 1992 (28.04.92)

(71) Applicant (for all designated States except US): HIGHTECH RECEPTOR AB [SE/SE]; c/o Active, Skeppsbron 2, S-

(72) Inventors; and
(75) Inventors/Applicants (for US only): BJÖRCK, Lars [SE/SE]; Kornvägen 40, S-240 17 Södra Sandby (SE). SJÖBRING, Ulf [SE/SE]; Lilla Sigridsgatan 1, S-223 50 Lund (SE).

(74) Agents: BERG, S., A. et al.; H. Albihns Patentbyrå AB, Box 3137, S-103 62 Stockholm (SE).

(81) Designated States: JP, US, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).

Published

With international search report.

(54) Title: PROTEIN L AND HYBRID PROTEINS THEREOF

(57) Abstract

(30) Priority data:

9201331-7

211 20 Malmö (SE).

The invention relates to sequences of protein L which bind to light chains of immunoglobulins. The invention also relates to hybrid proteins thereof which are able to bind to both light and heavy chains of immunoglobulin G, in particular protein LG. The invention also relates to DNA-sequences which code for the proteins, vectors which include such DNA-sequences, host cells which have been transformed with the vectors, methods for producing the proteins, reagent appliances for separation and identification of immunoglobulins, compositions and pharmaceutical compositions which contain the proteins.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

•	AT AU BB BE BF BG BJ BR CA CF CG CH C1	Australia Barbados Belgium Burkina Faso Bulgaria Benin Brazil Canada Central African Republic Congo Switzerland	FR GA GB GN GR HU IE IT JP KP	France Gabon United Kingdom Guinea Greece Hungary Ireland Italy Japan Democratic People's Republic of Korea Republic of Korea Kazakhstan Lineburstein		MR MW NL NO NZ PL PT RO RU SD SE SK SN SU	Mauritania Malawi Netherlands Norway New Zealand Poland Portugal Romania Russian Federation Sudan Sweden Slovak Republic Senegal Soviet Union
	BJ	Benin					
	BJ	Benin					
	BR			-	`	RU	Russian Federation
	CA					SD	Sudan
	CF	Central African Republic	KP			_	
	CG	Cango					
	CH	Switzerland				_	
	Cl	Côte d'Ivoire					
	CM	Cameroon	1.1	Liechtenstein		TD .	Chad-
	CS	Czechoslovakia	LK	Sri Lanka			Togo
	CZ	Czech Republic	I.U -	Luxembourg	-	TC	
	DE	Germany	MC	Монасо		UA	Ukraine
	DK	Denmark	MÇ	Madagascar		US	United States of America
			MI.	Mali		VN	Vict Nam
	ES	Spain Finland	MN	Mongolia			•
	14 L	7-iniana					

10

Protein L and Hybrid Proteins Thereof

The present invention relates to sequences of protein L which bind to light chains of immunoglobulins. The invention also relates to hybrid proteins of protein L having the ability to bind to light chains of all Ig and also to bind to light and heavy chains of immunoglobulin G, DNA-sequences which code for the proteins vectors that contain such DNA-sequences, host cells transformed by the vectors, methods for preparing the proteins, reagent apparatus for separating and identifying immunoglobulins, compositions and pharmaceutical compositions which contain the proteins.

The invention relates in particular to the DNA-sequence and to the amino acid sequence of the light-chain forming domains of protein L.

Proteins which bind to the constant domains (of high affinity) of the immunoglobulins (Ig) are known. Thus, 20 protein A (from Staphylococcus aureus) (Forsgren, A. and Sjöquist, J. (1966) Protein A from Staphylococcus aureus. I. Pseudo-immune reaction with human gammaglobulin. J. Immunol. 97: 822-827) binds to IgG from various mammal species. The binding of protein A to IgG 25 is mediated essentially via surfaces in the Fc-fragment of the heavy chain of the IgG-molecule, although a certain bond is also effected with surfaces in the Fabfragment of the IgG. Protein A lacks the ability of binding to human IgG3 and neither will it bind to IgG 30 from several other animal species, such as important laboratory animals, for instance rats and goats, which limits the use of protein A.

Protein G (Björck, L. and Kronvall, G. (1984) Purification and some properties of streptococcal protein G, a

30

novel IgG-binding reagent. J. Immunol. 133: 969-974; Reis, K., Ayoub, E. and Boyl, M. (1984) Streptococcal Fc receptors. I. Isolation and partial characterization of the receptor from a group C streptococcus. J. Immunol. 132: 3091-3097) binds to heavy chains in human IgG and to all four of its subclasses and also to IgG from most mammals, including rats and goats.

Protein H (Åkesson, P., Cooney, J., Kishimoto, F. and
Björck, L. (1990) Protein H - a novel IgG binding bacterial protein. Molec. Immun. 27: 523-531) binds to the
Fc-fragment in IgG from human beings, monkeys and rabbits. However, the bond is weaker than in the case of protein G and A, which may be beneficial when wishing to break the bond with a weak agent, for instance when purifying proteins which are readily denatured with the aid of antibodies.

Protein M (Applicant's Patent Application PCT/SE 91100447) binds to the Fc-fragment in IgG from humans, monkeys, rabbits, goats, mice and pigs.

Protein L (Björck, L. (1988) Protein L, a novel bacterial cell wall protein with affinity to Ig L chains. J. Immunol. 140: 1194-1197), which binds to the light chains in immunoglobulins from all of the classes G, A, M, D and E is known (USP 4,876,194). The amino acid sequence and the binding domains of this protein, however, have hitherto been unknown.

The aforesaid proteins can be used in the analysis, purification and preparation of antibodies and for diagnostic and biological research.

35 The elimination of immunoglobulins, with the aid of

plasmapheresis, can have a favourable effect on some autoimmune diseases. A broadly binding protein would be an advantage when wishing to eliminate all classes of antibodies in this context.

5

10

15

20

25

30

35

It has long been known that infectious conditions can be prevented or cured with the introduction of an immune serum, i.e. a serum which is rich in antibodies against the organism concerned or its potentially harmful product. Examples hereof are epidemic jaundice, tetanus, diphtheria, rabies and generalized shingles. Antibodies against a toxic product may also be effective in the case of non-infectious occasioned conditions. Serum produced in animals against different snake venoms is the most common application in this respect. However, the administration of sera or antibody preparations is not totally without risk. Serious immunological reactions can occur in some cases. Singular cases of the transmission of contagious diseases, such as HIV and hepatitis through the agency of these products have also been described. In order to avoid these secondary effects, it has been desirable to produce therapeutic antibodies in test tubes. A large number of novel techniques for the preparation of antibodies in test tubes have been proposed in recent years. Examples of such techniques are hybridom techniques, synthesis of chimaantibodies and the preparation of antibodies in bacteria. These techniques also enable antibodies to be specially designed which can further widen the use of such molecules as therapeutics, for instance in the case of certain tumour-diseases. In the case of some of these novel methods, however, the product totally lacks the Fc-fragment to which all of the described IgG-binding proteins, with the exception of protein L, bind. There is consequently a need of a process for purifying anti-

10

15

20

25

30

35

Δ

bodies for therapeutic use, wherein proteins which have a broad binding activity/specificity, can be of value.

It has long been possible to utilize the antibody reaction with its high grade specificity for diagnosing past or, in some cases, ongoing infections with different parasites. This indirect method of indicating infectious agents is called serology and, in many cases, may be the only diagnostic alternative. In certain cases, it can also be of interest to exhibit specific IgE- or IgAantibodies. When diagnosing with the aid of serology, the antigen is most often fastened to a solid phase, whereafter serum taken from the patient is incubated with the antigen. Antibodies that have been bound from the patient can then be detected in different ways, often with the aid of a secondary antibody (for instance, an antibody which is directed against the light chains of human antibodies) to which an identifiable label has been attached, such as alkaline phosphatase, biotin, radioactive isotopes, fluorescein, etc. In this context, a protein having a broad Ig binding capacity can be used as an alternative to secondary antibodies.

There are a number of non-therapeutic and non-diagnostic reasons for the necessity to bind antibodies. Antibodies are often used in research, both for detection and for purifying the antigen against which they are directed. All techniques which facilitate the purification of antibodies and, in particular, techniques which enable different classes to be purified, are of interest in this context.

Consequently, there is a serious need of a protein which has a broad binding activity/specificity and which binds to several different classes of immunoglobulins from different animal species. At present, there is no known

10

15

20

25

30

35

protein which will bind to all immunoglobulin classes. The earlier known proteins A, G, H and M bind only to heavy chains in IgG. The known protein L (Björck et al, 1988) binds to the light χ -chains and γ -chains in immunoglobulins of all classes, although the bonds are much weaker on the κ -chains. Applicant has charted protein L, has determined the amino acid sequence for protein L, has identified the light-chain binding domains on protein L, and has used these to produce hybrid proteins which possess the IgG-Fc-binding domains of protein G. The Applicant is able to show through protein LG that a protein of broader binding activity/ specificity can be produced thereby. The aforesaid proteins A, G, H and M bind to the same surfaces, or to very closely lying surfaces on IgG-Fc. The protein L which binds to light chains can thus be combined with any other functionally similar protein which binds to the Fc-fragment of heavy chains. A similar broadening of the Ig-binding activity is achieved with all alternatives.

Thus, the present invention relates to the sequence of protein L which binds to light chains in Ig and has the amino acid sequence disclosed in Figure 1, and variants, subfragments, multiples or mixtures of the domains B1-B5 having the same binding properties. The invention also relates to a DNA-sequence which codes for such protein sequences, for instance the DNA-sequence in Figure 1.

The invention is concerned with a hybrid protein which is characterized by comprising domains which bind to the light χ -chains and λ -chains in immunoglobulins of all classes, and also comprises domains which bind to heavy chains in immunoglobulin G, wherein those domains which bind to the light chains are chosen from among the B1-, B2-, B3-, B4- and B5-domains in protein L (see Claim 1)

25

30

35

and those domains which bind to heavy chains of immunoglobulins are chosen from th C1-, C2- and C3-domains in protein G; the A-, B- and C1-domains from protein H; the A-, B1-, B2- and S-domains in protein M1 or the E-, D-, A-, B- and C-domains in protein A (see Figure 6) and variants, subfragments, multiples or mixtures of these domains that have the same binding properties which bind to heavy chains of immunoglobulins.

By subfragment is meant a part-fragment of the given 10 domains or fragments which include parts from the various domains having mutually the same binding properties. By variants is meant proteins or peptides in which the original amino acid sequence has been modified or changed by insertion, addition, substitution, inversion 15 or exclusion of one or more amino acids, although while retaining or improving the binding properties. The invention also relates to those proteins which contain several arrays (multiples) of the binding domains or mixtures of the binding domains with retained binding 20 properties. The invention also relates to mixtures of the various domains of amino acid sequences having mutually the same binding properties.

The invention relates in particular to a hybrid protein designated LG, and is characterized in that the hybrid protein includes the B-domains in protein L which bind to the light chains in immunoglobulins, and the C1-domains and C2-domains in protein G which bind to heavy chains and have the amino acid sequence disclosed in Figure 3. The invention also relates to variants, subfragments, multiples or mixtures of these domains.

Protein LG is a hybrid protein having a molecular weight of about 50 kDa (432 amino acids) and comprising four domains, each of which binds to light chains in immuno-

10

15

20

25

30

35

globulins, and two IgG-binding domains from protein G. The hybrid protein combines a br ad IgG-binding activity, deriving from the high-grade binding ability of protein G to the Fc-fragment of the heavy chain on IgG with the ability of the protein L to bind to light chains of all classes of immunoglobulins. Thus, protein LG binds polyclonal human IgG, IgM, IgA, IgD and IgE. The affinity for human polyclonal IgG is 2 x 10^{10} M⁻¹. All four human immunoglobulin classes are bound. Binding to human IgG is effected with both the κ -and the λ chain. Both the Fc-fragment and the Fab-fragment of IgG are bound to the hybrid protein. The protein also binds human IgA-, IgD-, IgE- and IgM-antibodies. The bond is stronger to human immunoglobulins which carry χ than to those which carry the λ -isotope of light chains. IgG from most mammals will be bound by protein LG, thus also IgG from goats and cows, which do not bind to protein L. However, rabbit-IgG which binds relatively weakly to protein L will bind well to the fusion protein. IgM and IgA-antibodies from mice, rats and rabbits will be bound to the protein.

Protein LG is highly soluble. It is able to withstand heat and will retain its binding properties even at high temperatures. The binding properties also remain in a broad pH-range of 3-10. The protein withstands detergent and binds marked or labelled proteins subsequent to separation in SDS-PAGE and transference to membranes with elektroblotting. The protein can be immobilized on a solid phase (nitrocellulose, Immobilon®, polyacrylamide, plastic, metal and paper) without losing its binding capacity. The binding properties are not influenced by marking with radioactive substances, biotin or alkaline phosphatase. (The binding abilities of the protein LG are disclosed in Example 3).

10

15

The protein comprises 432 amino acids and has a molecular weight of 50 kDa deriving therefrom. The sequence is constructed of an ala sequence of the three last amino acids in the A-domain of the protein L (val-glu-asn), this ala sequence being unrelated to the two proteins, whereafter the four mutually high-grade homologous Bdomains from protein L follow. The first of the B-domains is comprised of 76 amino acids, and the remaining domains are each comprised of 72 amino acids. The first nine amino acids from the fifth B-domain are included and followed by two non-related amino acids (pro-met). The protein G-sequences then follow. The last amino acid in the so-called S-domain from protein G is followed by an IgG-binding domain from protein G (C1; 55 amino acids), the intermediate D-region (15 amino acids) and the second IgG-binding C-domain (C2; 55 amino acids). The last amino acid is a methionine, which occurs in natural protein G as the first amino acid in the so-called W-region.

20

25

30

35

The invention also relates to DNA-sequences which code for the aforesaid proteins.

The gene which codes for the IgG-binding amino acid sequences can be isolated from the chromosomal DNA from Staphylococcus aureus based on the information on the DNA-sequence for protein A (S. Löfdahl, B. Guss, M. Uhlen, L. Philipsson and M. Lindberg. 1983. Gene for staphylococcal protein A. Proc. Natl. Acad. Sci. USA. 80: 697-701) and Figure 6, or from G-streptococcus, preferably strain G 148 or C-streptococcus, preferably strain Streptococcus equisimilis C 40, based on the information on protein G (B. Guss, M. Eliasson, A. Olsson, M. Uhlen, A.-K. Frej, H. Jörvall, I. Flock and M. Lindberg. 1986. Structure of the IgG-binding

10

15

20

25

30

35

regions of streptococcal protein G. EMBO. J. 5: 1567-1575) and Figure 6, or from group A-streptococcus, e.g. S. pyogenes (type M1) based on the information on the DNA-sequence for protein H (H. Gomi, T. Hozumi, S. Hattori, C. Tagawa, F. Kishimoto and L. Björck. 1990. The gene sequence and some properties of protein H - a novel IgG binding protein J. Immunol. 144: 4046-4052) and Figure 6, or from the chromosomal DNA in group A-streptococcus type M1 based on the information on the DNA-sequence for protein M (Applicant's Patent Application, PCT/SE 91100447) and Figures 6 and 7. The gene which codes for the protein that binds to light chains can be isolated from the chromosomal DNA from Pepto-coccus magnus 312 based on the information on the DNA-sequence for protein L in Claim 2.

By using the chromosomal DNA't obtained from the aforesaid bacteria as a template, a DNA-fragment defined with the aid of two synthetic oligonucleotides can then be specifically amplified with the aid of PCR (Polymerase Chain Reaction). This method also enables recognition sites to be incorporated for restriction enzymes in the ends of the amplified fragments (PCR technology, Ed: PCR Technology. Principles and Applications for DNA Amplification. Ed. Henry Erlich. Stockton Press, New York, 1989). The choice of recognition sequences can be adapted in accordance with the vector chosen to express the fragment or the DNA-fragment or other DNA-fragments with which the amplified fragment is intended to be combined. The amplified fragment is then cleaved with the restriction enzyme or enzymes concerned and is combined with the fragment/the other fragments concerned and the fragments are then cloned together in the chosen vector (in this case, the expression vector) (Sambrook, J.E. Fritsch and T. Maniatis, 1989, Molecular cloning: A laboratory manual, 2nd Ed. Cold Spring Harbor Laborato-

ries, Cold Spring Harbor, New York, USA). The plasmid vector pHD313 can be used (Dalböge, H.E. Bech Jensen, H. Töttrup, A. Grubb, M. Abrahamson, I. Olafsson and S. Carlsen, 1989. High-level expression of active human cystatin C in Escherichia coli. Gene, 79: 325-332), alternatively one of the vectors in the so-called PET-series (PET 20, 21, 22, 23) retailed by Novagen (Madison, Wisconsin, USA).

- The hybrid proteins are then incorporated in an appropriate host, preferably E. coli. The invention also relates to such hosts as those in which the hybrid proteins are incorporated.
- Those clones which produce the desired proteins can be selected from the resultant transformants with the aid of a known method (Fahnestock et al., J. Bacteriol. 167, 870 (1986).
- When the proteins that can bind to the light chains in the immunoglobulins and to the heavy chains in IgG have been purified from the resultant positive clones with the aid of conventional methods, the binding specificities of the proteins are determined for selection of those clones which produce a protein that will bind to the light chains in immunoglobulins and to the heavy chains in IgG.
- Subsequent to having isolated plasmid DNA't in said clone with conventional methods, the DNA-sequence in the inserted material is determined with known methods (Sanger et al., Proc. Natl. Acad. Sci. USA 74, 5463 (1977).
- The invention also relates to DNA-sequences which hybridize with said identified DNA-sequences under conven-

tional conditions and which code for a protein that possesses the desired binding properties. Strict hybridizing conditions are preferred.

Expression of the genes can be effected with expression vectors which have the requisite expression control regions, the structural gene being introduced after said regions. As illustrated in Figure 1 and Claim 2, the structural gene can be used for protein LG or other hybrid proteins with protein L.

With regard to expression vectors, different host-vector-systems have been developed, of which the most suitable host-vector-systems can be selected for expression of the genes according to the present invention.

The present invention also relates to a method of producing the inventive hybrid proteins by cultivating a host cell which is transformed with an expression vector in which DNA't which codes for the proteins according to the invention is inserted.

This method includes the steps of

- 25 (1) inserting into a vector a DNA-fragment which codes for the hybrid proteins;
 - (2) transforming the resultant vector into an appropriate host cell;
 - (3) cultivating the resultant, transformed cell for preparation of the desired hybrid protein; and
 - (4) extracting the protein from the culture.

35

30

15

20

In the first step, the DNA-fragment which codes for the hybrid protein is inserted in a vector which is suitable for the host that is to be used to express the hybrid protein. The gene can be inserted by cleaving the vector with an appropriate restriction enzyme, and then legating the gene with the vector.

In the second step, the vector with the hybrid plasmid is inserted into host cells. The host cells may be Escherichia, coli, Bacillus subtilis or Saccharomyces cerevisiae or other suitable cells. Transformation of the expressions hybrid vector into the host cell can be effected in a conventional manner and clones which have been transformed can then be selected.

15

5

10

In the third step, the obtained transformants are cultivated in an appropriate medium for preparation of the desired proteins by expression of the gene coded for the hybrid protein.

20

25

30

In the fourth step, the desired protein is extracted from the culture and then purified. This can be achieved with the aid of known methods. For instance, the cells can be lysed with the aid of known methods, by treating the cells with ultrasonic sound, enzymes or by mechanical degradation. The protein which is released from the cells or which excretes in the medium can be recovered and purified with the aid of conventional methods often applied within the biochemical field, such as ion-exchange chromatography, gel filtration, affinity chromatography with the use of immunoglobulins as ligands, hydrophobic chromatography or reverse-phase chromatography. These methods can be applied individually or in suitable combinations.

35

As before mentioned, the inventive proteins may be used for binding, identifying or purifying immunoglobulins. They can also be bound to pharmaceuticals and used in formulations which have delayed release properties. To this end, the protein may be present in a reagent appliance for pharmaceutical composition in combination with appropriate reagents, additives or carriers.

in a PBS-solution (phosphate-buffered physiological salt solution) pH 7.2 with 0.02% NaN3. It can also be used connected to a solid phase, such as carbohydrate-based phases, for instance CNBr-activated sephanose, agarose, plastic surfaces, polyacrylamide, nylon, paper, magnetic spheres, filter, films. The proteins may be marked with biotin, alkaline phosphatase, radioactive isotopes, fluorescein and other fluorescent substances, gold particles, ferritin, and substances which enable luminescence to be measured.

20

25

30

35

Other proteins may also be used as carriers. These carriers may be bound to or incorporated in the proteins, in accordance with the invention. For instance, it is conceivable to consider the whole of proteins A, G, H, M as carriers for inserted sequences of protein L which bind to light chains. In turn, these carriers can be bound to the aforesaid carriers.

The pharmaceutical additions that can be used are those which are normally used within this field, such as pharmaceutical qualities of mannitol, lactose, starch, magnesium stearate, sodium saccharate, talcum, cellulose, glycose, gelatine, saccharose, magnesium carbonate and similar extenders, such as lactose, dicalcium phosphate and the like; bursting substances, such as starch or derivatives thereof; lubricants such as magnesium

20

25

30

stearate and the like; binders, such as starch, gum aribicum, polyvinylpyrrolidone, gelatine, cellulose and derivatives thereof, and the like.

The invention will now be described in more detail with reference to the accompany drawings, in which

Figure 1 illustrates the plasmid pHD389; the ribosomal binding sequence, the sequence for the signal peptide from ompA and recognition sequence for several restriction enzymes are shown;

Figure 2 illustrates the amino acid and nucleic acid sequence for protein LG;

Figure 3 is a schematic overall view of the production of protein L;

Figure 4 is a schematic overall view of the production of protein LG;

Figures 5a, 5b and 5c are schematic overall views of the production of the hybrid proteins LA, LM and LH respectively;

Figure 6 is a schematic inclusive illustration of protein A, G, H and M1. IgGFc-binding domains are for protein A: E, D, A, B and C; for protein G: C1, C2 and C3; for protein H: A and/or B; and for protein M1: A, B1, B2, B3 and S;

Figure 7 illustrates the amino acid and nucleic acid sequence for protein M1;

Figure 8 illustrates Western Blot for protein G, L and LG with certain immunoglobulins and immunoglubulin fragments; and

5 Figure 9 illustrates Slot-Blot for protein L, G and LG with IgG, Ig χ and Ig Fc.

The amino acid and nucleic acid sequence of the lightchain binding domains of protein L is illustrated in Claims 1 and 2 respectively.

It will be observed that the drawings are not to scale.

Example 1

15

20

25

10

Cloning and expression of the IgG-light-chain-binding domains in Protein L

Construction of synthetic oligonucleotides (primers) for amplifying sequences coded for protein L, domain B1-B4

It has been found that a protein L peptide (expressed in E. coli) constructed of the sequence ala-val-glu-asn-domain B1 (from protein L) binds to the light chains of the immunoglobulins (W. Kastern, U. Sjöbring and L. Björck. 1992. Structure of peptostreptococcal protein L and identification of a repeated immunoglobulin light chain-binding domain. J. Biol. Chem. in-print). Since this simple protein L-domain has a relatively low affinity to Ig, $(1 \times 10^7 \text{ M}^{-1})$, and since the naturally occurring protein L which is constructed of several mutually similar domains (B1-B5) has a high affinity to Ig $(1 \times 10^{10} \text{ M}^{-1})$ four of these domains have been expressed together in the following way:

30

PL-N and PL-C1 are synthetic olig nucleotides (manufactured by the Biomolecular Unit at Lund University (Sweden) in accordance with Applicant's instructions) which have been used to amplify a clonable gene fragment which is amplified with PCR (Polymerase Chain Reaction) and which codes for four Ig-binding protein L domains (alaval-glu-asn-B1-B2-B3-B4-lys-lys-val-asp-glu-lys-pro-glu-glu). Amino acids in the protein L-sequence are given for the primer which corresponds to the coded strand (PL-N):

PL-N: 5'-GCTCAGGCGCGCCGGTAGAAAATAAAGAAGAAACACCAGAAAC-3' valgluasnlysglugluthrproglu

5'-end of this oligonucleotide is homologous with the coded strand in the protein L-gene (emphasized): those codons which code for the last three amino acids in the A-domain (val-glu-asn) are followed by the codons for the first six amino acids in the first of the Ig-binding domains in protein L (B1).

PL-C1: 5'-CAGCAGCAGGATTCTTATTATTATTCTTCTGGTTTTTCGTCAACTTT
CTT-3'

25

10

15

20

This oligonucleotide is homologous with the opposing non-coding strand in the gene for protein L (the sequence corresponds to the first nine amino acids in domain B5).

30

35

DNA-fragments which have been amplified with the aid of PL-N contain the recognition sequence for the restriction enzyme **HpaII** (emphasized) immediately before the codon which is considered to code for the first amino acid (val) in the expressed protein L-fragment. The fragment which is cleaved with **HpaII** can be ligated with

25

30 -

35

DNA (in this case, consisting of the used expression vector pHD389) which has been cleaved with the restriction enzyme Narl. The DNA-fragment that has been cleaved with Hpall and ligated with vector pHD389, which has been cleaved with Narl, will be translated in the correct reading frame. The construction results in translation of an additional amino acid (ala) immediately in front of the first amino acid in protein L.

DNA-fragments which have been amplified with the aid of PL-C1 will contain the recognition sequence for the restriction enzyme BamHI (overlined above the sequence) immediately after the sequence which codes for the last amino acid in the expressed protein L-fragment (glu).

The vector pHD389 contains a unique recognition sequence for BamHI as part of its so-called multiple cloning sequence which follows the NarI recognition sequence.

DNA-fragments which have been amplified with the aid of PL-C1 will include two so-called stop-codons (emphasized) which results in translation of the fragment inserted in the vector to cease.

The sequence which was considered to be amplified contains no internal recognition sequences for the restriction enzymes HpaII or BamHI.

Amplifying and cloning procedures

(PCR) (Polymerase Chain Reaction) was effected with a protocol described by Saiki, R.D. Gelfand, S. Stoffel, S. Scharf, R. Higuchi, G. Horn, K. Mullis and H. Erlich, 1988; Primer-directed enzymatic amplification of DNA with a thermostable DNA polymerase. Science 239: 487-49127; PCR was effected in a Hybaid Intelligent Heating-block (Teddington, UK): 100 μl of a reaction mixture contained 50 mM KCl, 10 mM Tris-HCl, pH 8.3, 1.5 mM

10

15

20

25

30

35

 $MgCl_2$, 100 μ/ml gelatine, 300 μM with respect to each of the deoxynucleotides (dATP, dCTP, dGTP, dTTP), (Pharmacia), 20 pmol of each of the oligonucleotides PL-N and PL-C1, 10 μ l of a target (template) DNA-solution containing 0.1 mg/ml of chromosomal DNA from Peptostreptococcus magnus, species 312. The mixture was covered with mineral oil (Sigma) and DNA't was denatured by heating to 98°C for 10 minutes. 2.5 units of AmpliTag (Perkin Elmer Cetus, Norwalk, CT) were added and PCR was then carried out with 25 cycles consisting of a denaturing step at 94°C for 1 minute, followed by a hybridizing step at 56°C for 1 minute, and finally by an extension step at 72°C for 1 minute. Amplified DNA was analyzed by electrophoresis in agarose gel. The amplified DNA't was cleaved with the restriction enzymes HpaII (Promega), (8 units/ μ g amplified DNA) and BamHI (Promega), (10 units/ μ g amplified DNA) at 37°C. The thus amplified and subsequently cleaved DNA-product was isolated by electrophoresis in a 2% (weight by volume) agarose gel (NuSieve agarose, FMC Biproducts) in a TAE-buffer (40 Mm Tris, 20 Mm Na-acetate, 2 Mm EDTA, Ph 8.0). The resulting 930 base-pair large fragment was cut from the gel. The DNA concentration in this removed gel-piece was estimated to be 0.05 mg/ml. The agarose-piece containing the cleaved, amplified fragment was melted in a water bath at 65°C, whereafter the fragment was allowed to cool to 37°C. 10 μ l (0.5 μ g) of this DNA was transferred to a semimicrotube (Sarstedt) preheated to 37°C, whereafter 1 μ l of the vector pHD389 was immediately added and cleaved with NarI (Promega) and BamHI, 1 μ l 10xligas-buffer (Promega and 1 μ l T4 DNA-ligase (Promega; 1 unit/ μ l). The ligating reaction was then used to transform E. coli, strain LE392, which had been competent in accordance with the rubidium/calcium-chloridemethod as described by Kushner (1978). Molecular biological standard methods have been used in the manipulation

. 1

5

of DNA (Sambrook, J.E. Fritsch and T. Maniatis, 1989. Molecular cloning: A laboratory manual. 2nd Ed. Cold Spring Harbor Laboratories, Cold Spring Harbor, New York, USA). The cleaving and ligating conditions recommended by the manufacturer of DNA-ligase and restriction enzymes have been followed in other respects.

Expression system

The vector pHD389 (see Figure 2) is a modified variant 10 of the plasmid pHD313 (Dalböge, H.E. Bech Jensen, H. Töttrup, A. Grubb, M. Abrahamson, I. Olafsson and S. Carlsen, 1989. High-level expression of active human cystatin C in Escherichia coli. Gene, 79: 325-332). The vector, which is replicated in E. coli (contains ori = 15 origin of replication from plasmid pUC19) is constructed so that DNA-fragments which have been cloned into the cleaving site of Narl will be transcribed and translated downstream of and in the immediate vicinity of the signal peptide (21 amino acids), from envelope-protein 20 ompA from E. coli. Translation will be initiated from the codon ATG which codes for the first amino acid (methionine) in the signal peptide. This construction permits the translated peptide to be transported to the periplasmic space in E. coli. This is advantageous, 25 since it reduces the risk of degradation of the desired product of enzymes occurring intracellularly in E. coli. Moreover, it is easier to purify peptides which have been exported to the periplasic space. Unique recognition sequences (multiple cloning sequences) for sever-30 al other restriction enzymes, among them ecoRI, SalI and BamHI are found immediately after the NarI cleaving site. An optimized so-called Shine-Dalgarno-sequence (also called ribosomal binding site, RBS) is found seven nucleotides upstream from the ATG-codon in the signal 35 sequence from ompA, this optimized sequence binding to a

10

complementary sequence in 16S rRNA in the ribosomes and is responsible for the translation being initiated in the c rrect place. The transcription of such DNA as that which is co-transcribed with the signal sequence for ompA is controlled by the $P_R^{}$ -promotor from coliphage $\lambda.$ The vector also contained the gene for cI857 from coliphage λ whose product down-regulates transcription from P_p (and whose product is expressed constitutively). This $c\bar{1}857$ -mediated down-regulation of transcription from P_R is heat-sensitive. The transcription regulated from this promotor is terminated with the aid of a so-called rhoindependent transcription terminating sequence (forms a structure in DNA't which results in the DNA-dependent RNA-polymerase leaving the DNA-strand) which is placed in the vector immediately downstream of the multiple cloning sequence. The plasmid also carries the B-lactamase gene (from the plasmid pUC19) whose product permits ampicillin-selection of E. coli clones that have been transformed by the vector.

20

25

30

35

15

Selection of protein L-producing clones

The transformed bacteria are cultivated, or cultured, on culture plates with an LB-medium which also contained ampicillin in a concentration of $100~\mu g/ml$. Cultivation of the bacteria progressed overnight at $30\,^{\circ}$ C, whereafter the bacteria were transferred to an incubator where they were cultivated for a further 4 hours at $42\,^{\circ}$ C. The plates were kept in a refrigerator overnight. On the next day, the colonies were transferred to nitrocellulose filters. Filters and culture plates were marked so as to enable the transferred colonies to be readily identified on respective culture plates. The culture plates were again incubated overnight at $30\,^{\circ}$ C, so that remaining rests of transferred bacteria colonies could again gr w. The plates were then kept in a refrigerator.

15

20

25

30

35

The bacteria in the colonies on the nitr celluloseimpressions were lysed by incubating the filter in 10% SDS for 10 minutes. Filters c ntaining lysed bacteria were then rinsed with a blocking buffer which comprised PBS (pH 7.2) with 0.25% gelatine and 0.25% Tween-20 (four baths, 250 ml each at 37°C), whereafter the filter was incubated with radioactively marked (marked with 125 I in accordance with the chloramin-T-method) Ig- κ chains (20 ng/ml in PBS with 0.1% gelatine). The incubation took place at room temperature over a period of 3 hours, whereafter non-bound radioactively marked protein was rinsed-off with PBS (pH 7.2) containing 0.5 M NaCl, 0.25% gelatine and 0.25% Tween-20 (four baths, 250 ml each at room temperature). All filters were exposed to X-ray film. Positive colonies were identified on the original culture plate. Clones which reacted with Ig-x-chains were selected and analyzed with respect to the size on the DNA-fragment introduced in the vector. One of these clones was selected for the production of protein L, pHDL. The DNA't introduced from this clone into plasmid pHD389 was sequenced. The DNAsequence was found to be in full agreement with corresponding sequences (B1-B4 and 21 bases in B5) in the gene for protein L from Peptostreptococcus magnus; strain 312. The size and binding properties of the protein produced by clone pHDL was analyzed with the aid of SDS-PAGE (see Figure 8), dot-blot experiment (see Figure 9) and competitive binding experiments.

Production of protein L

Several colonies from a culture plate with <u>E. coli</u> pHDL were used to inoculate a preculture (LB-medium with an addition of 100 mg/l ampicillin), which was cultured at 28°C overnight. On the following morning, the preculture was transferred t a larger volume (100 times the volume

10

15

20

25

of the preculture) of fresh LB-medium containing ampicillin (100 mg/l) and was cultured in shake-flasks (200 rpm), (or fermentors) at 28°C. The culture temperature was raised to 40°C (induction of transcription) when the absorbency value at 620 nm reached 0.5. Cultivation then continued for 4 hours (applied solely to cultivation in shake-flasks). Upon completion of the cultivation process, the bacteria were centrifuged down. The bacteria were then lysed with an osmotic shock method at 4°C (Dalböge et al., 1989 supra). The lysate was adjusted to a pH = 7. Remaining bacteria rests were then centrifuged down, whereafter the supernatent was purified on IgGsepharose in accordance with earlier described protocol for protein G and protein L (U. Sjöbring, L. Björck and W. Kastern. 1991. Streptococcal protein G: Gene structure and protein binding properties. J. Biol. Chem. 266: 399-405; W. Kastern, U. Sjöbring and L. Björck. 1992. Structure of peptostreptococcal protein L and identification of a repeated immunoglobulin light chain-binding doman. J. Biol. Chem. in-print.

The expression system gave about 20 mg/l of protein L when cultivation in shake-flasks. The culture was deposited at DSSM, Identification Reference DSSM $\underline{E.\ coli}$ LE392/pHDL.

Example 2

Cloning and expression of protein LG

5 Construction of oligonucleotides (primers) for amplifying sequences which code for protein LG

Protein L

It has been found that a protein L-peptide (expressed in E. coli) constructed of the sequence ala-val-glu-asn-domain B1 (from protein L) will bind to the light chains of the immunoglobulins (Kastern, Sjöbring and Björck, 1992, J. Biol. Chem. in-print). Since the affinity of this simple domain to Ig is relatively low (1 x 10⁻⁷ M⁻¹) and since the naturally occurring protein L, which is comprised of several mutually similar domains (B1-B5) has a higher affinity to Ig (1 x 10¹⁰ M⁻¹), four of these domains have been expressed together in the following way:

PL-N and PL-C2 are synthetic oligonucleotides (manufactured at the Biomolecular Unit at Lund University (Sweden) in accordance with Applicant's instructions) which were used, with the aid of PCR (Polymerase Chain Reaction) to amplify a clonable gene fragment, called B1-4, which codes for four Ig-binding protein L domains (alaval-glu-asn-B1-B2-B3-B4-lys-lys-val-asp-glu-lys-pro-glu-glu):

30

25

PL-N: 5'-GCTCAGGCGGCGCCGCTAGAAAATAAAGAAGAAACACCAGAAAC-3' valgluasnlysglugluthrproglu

P1-C2: 5'-CAGCAGCAGCCATGGGTTCTTCTGGTTTTTCGTCAACTTTCTTA-

35

3'

10

Amino acids have been shown under corresponding triplets in the coded strand. DNA-fragments which have been amplified with the aid of PL-N contain the recognition sequence for the restriction enzyme HpaII immediately upstream of the triplet which codes for the first amino acid (val) in the expressed protein L-fragment. The fragment that has been cleaved with HpaII can be ligated with DNA (in this case, the used expression vector pHD389) which has been cleaved with Narl. The construction results in translation of an extra amino acid (ala) immediately upstream of the first amino acid in the protein L-fragment. The DNA-fragment that has been amplified with the aid of PL-C2 will contain the recognition sequence for the restriction enzyme Ncol (emphasized) immediately downstream of the sequence which codes for the last amino acid in the expressed protein L-fragment (glu). Amplified fragments which have been cleaved with Ncol can be ligated to the Ncol-cleaved, PCR-generated protein-asp-CDC-met-fragment (see below).

20

25

30

35

15

Protein G

It is known that a simple C-domain from protein G will bind to IgG (B. Guss, M. Eliasson, A. Olsson, M. Uhlen, A.-K. Frej, H. Jörnvall, I. Flock and M. Lindberg. 1986. Structure of the IgG-binding regions of streptococcal protein G. EMBO. J. 5: 1567-1575). The strength at which a simple C-domain binds to IgG is relatively low $(5 \times 10^7 \text{ M}^{-1})$. A fragment which consists of two C-domains with an intermediate D-region having a length of 15 amino acids, however, has a considerably higher affinity to IgG (1 x 10 9 M $^{-1}$). CDC-N and CDC-C are oligonucleotides which have been used as PCR-primers to amplify a clonable DNA-fragment, designated CDC, which

codes for two IgG-binding protein G-domains (pro-met-asp-CDC-met).

CDC-N: GGCCATGGACACTTACAAATTAATCCTTAATGGT metaspthrtyrlysleuileleuasngly

5

35

CDC-C: CAGGTCGACTTATTACATTTCAGTTACCGTAAAGGTCTTAGT

Amino acids in the resultant sequence have been shown beneath the primer of the coding strand. DNA-fragments 10 which have been amplified with the aid of CDC-N contain the recognition sequence for the restriction enzyme Ncol (marked with a line above the sequence). Cleaved amplified fragments can be ligated with the fragment that has been amplified with the aid of PL-C2 and then cleaved 15 with Ncol. The fragment will therewith be translated to the correct reading frame. DNA-fragments which have been amplified with the aid of CDC-C will contain two socalled stop condons (emphasized) which terminate translation. The recognition sequence for the restriction 20 enzyme sall (marked with a line above the sequence) follows immediately afterwards, this sequence also being found in the expression vector pHD389 (see Figure 1).

Those sequences which code for the binding properties of protein L (B1-B5) and for protein G (CDC) respectively contain no internal recognition sequences for the restriction enzymes HpaII, SalI or NcoI.

30 Amplifification and cloning procedures

PCR (Polymerase Chain Reaction) was carried out in accordance with a protocol described by Saiki et al., 1988; PCR was carried out in a Hybaid Intelligent Heating-block (Teddington, UK): 100 μ l of the reaction mixture contained 50 mM KC1, 10 mM Tris-HCl, pH 8.3, 1.5

10

15

20

25

30

35

mM MgCl₂, 100 μ g/ml gelatine, 300 μ M with respect to each of the deoxynucleotides (dATP, dCTP, dGTP, dTTP), (Pharmacia). In order to amplify sequences which code for the light-chain binding parts of protein L, there were added 20 pmol of each of the oligonucleotides PL-N and PL-C2, and 10 μ l of a DNA-solution which contained 0.1 mg/ml of chromosomal DNA from Peptostreptococcus magnus, strain 312. By way of an alternative, 20 pmol were added to each of the oligonucleotide pairs CDC-N and CDC-C and 10 μ l of a DNA-solution which contained 0.1 mg/ml of chromosomal DNA from a group C streptococcus strain (Streptococcus equisimilis) called C40 (U. Sjöbring, L. Björck and W. Kastern. 1991. Streptococcal protein G: Gene structure and protein binding properties. J. Biol. Chem. 266: 399-405 or with Ncol and SalI (10 U/ μ g PCR-product), (for CDC) at 37°C. The thus amplified and subsequently cleaved DNA-fragments were then separated by electrophoresis in a 2% (weight by volume) agrose gel (NuSieve agarose, FMC Bioproducts) in a TAE-buffer (40 mM Tris, 20 mM aNa-cetate, 2 mM EDTA, pH 8.0). The resultant fragments, 930 bp (for B1-4) and 390 bp (for CDC) were cut from the gel. The concentration of DNA in the thus separated gel pieces was estimated to be 0.05 mg/ml. The agarose pieces cut from the gel and containing the cleaved, amplified fragments (B1-4 and CDC) were melted in a water bath at 65°C, whereafter they were allowed to cool to 37°C. 10 μ l (0.5 μ g) of this DNA were transferred to a semimicrotube (Sarstedt), preheated to 37°C, whereafter 1 μ l of the vector pHD389 which had been cleaved with NarI and Sall were added. 1 μ l 10 x ligase buffer (Promega) and 1 μ l T4 DNA-ligase (1 unit/ μ l) were also added. The ligating reaction was permitted to take place at 37°C for 6 hours. The cleaving and ligating conditions recommended by the producer of DNA-ligase and restriction enzymes (Promega) were followed in other respects. The

ligating reaction was then used to transform <u>E. coli</u>, strain LE392, which had been made competent in accordance with the rubidium-chloride/calcium-dichloride method as described by Kushner (1978). Manipulation of DNA was effected in accordance with molecular biological standard methods (Sambrook et al., 1989).

Expression system

The vector pHD389 (see Figure 2) is a modified variant 10 of the plasmid pHD313 (Dalböge et al., 1989). The vector which was replicated in E. coli (contains origin of replication from plasmid pUC19) is constructed such that DNA-fragments which have been cloned in the cleaving site for Narl will be expressed immediately after, or 15 downstream, of the signal peptide (21 amino acids) from the envelope protein ompA from E. coli. Translation will be initiated from the ATG-codon which codes for the first amino acid (methionine) in the signal peptide. The construction with an E. coli-individual signal sequence 20 which precedes the desired peptide enables the translated peptide to be transported to the periplasmic space in E. coli. This is beneficial since it reduces the risk of degradation of the desired product through the intracellular occurrent enzymes of E. coli. Furthermore, it is 25 easier to purify peptides which have been exported to the periplasmatic space. Unique recognition sequences (multiple cloning sequences) for several other restriction enzymes, among them EcoRI, SalI and BamHI are present immediately downstream of the NarI cleaving 30 site. An optimized so-called Shine-Dalgarno sequence (also called ribosomal binding site, RBS) is found seven nucleotides upstream of the ATG-codon in the signal sequence from ompA, this optimized Shbine-Dalgarno sequence binding to a complementary sequence in 16S rRNA 35 in the ribosomes and in a manner to decide that the

translati n is initiated in the c rrect place. The transcription of such DNA as that which is co-transcribed with the signal sequence for mpA is controlled by the P_{R} -promotor from coliphage λ . The vector also contains the gene for cI857 from coliphage λ , the product of which regulates-down transcription from $P_{_{\rm D}}$ and the product of which is expressed constitutively. This cI857-mediated down-regulation of transcription from Pp is heat-sensitive. Transcription which is regulated, or controlled, from this promotor will be terminated with the aid of a so-called rho-independent transcription terminating sequence which is inserted in the vector immediately downstream of the multiple cloning site. The plasmid also carries the gene for 8-lactamase (from the plasmid pUC19), the product of which permits ampicillinselection of E. coli clones that have been transformed with the vector.

Selection of protein LG-produced clones

20

25

30

35

10

15

The transformed bacteria are cultivated on culture plates with LB-medium which also contained ampicillin in a concentration of 100 μ g/ml. The bacteria were cultivated overnight at 30°C, whereafter they were transferred to a cultivation cabinet (42°C) and cultured for a further four (4) hours. The plates were stored in a refrigerator overnight. On the following day, the colonies were transferred to nitrocellulose filters. The filters and culture plates were marked, so that the transferred colonies could later be identified on the culture plate. The culture plates were again incubated overnight at 30°C, so that rests of transferred bacteria colonies remaining on the plates could again grow. The plates were then stored in a refrigerator. The filter was incubated in 10% SDS for 10 minutes, so as to lyse the bacteria in the c lonies on the nitrocellulose

10

15

20

25

30

35

impression. Filters containing lysed bacteria were then rinsed with a blocking buffer c nsisting of PBS (pH 7.2) with 0.25% gelatine and 0.25% Tween-20 (four baths of 250 ml at 37°C), whereafter the filter was incubated with radioactively (marked with ^{125}I according to the chloromine-T-method) marked Ig-x-chains (20 ng/ml) in PBS with 0.1% gelatine). The incubation process took place at room temperature for four (4) hours, whereafter non-bound radioactively marked protein was rinsed-off with PBS (pH 7.2) containing 0.5 M NaCl, 0.25% gelatine and 0.25% Tween-20 (four baths, 250 ml each at room temperature). All filters were exposed to X-ray film. Positive colonies on the original culture plate were identified. A number of positive colonies were recultivated on new plates and new colony-blot experiments were carried out with these plates as a starting material with the intention of identifying E. coli colonies which bind IgG Fc. These tests were carried out in precisely the same manner as that described above with respect to the identification of E. coli-colonies which expressed Ig light-chain-binding protein, with the exception that a radioactively marked (125 I) IgG Fc (20 ng/ml) was used as a probe. Clones which reacted with both proteins were selected and analyzed with regard to the size of the DNA-fragment introduced in the vector. One of these clones was chosen for production of protein LG, pHDLG. The DNA't taken from this clone and introduced into plasmid pHD389 was sequenced. The DNA-sequence exhibited full agreement with corresponding sequences (B1-B4 and 21 bases in B5) in the gene for protein L from Peptostreptococcus magnus, strain 312, and with C1DC2 sequence in group C streptococcus strain C40. The size and binding properties of the protein produced from clone pHDLG was analyzed with the aid of SDS-PAGE (see Figure 8), dot-blot experiment (see Figure 10) and competitive binding experiments.

10

15

20

35

Pr ducti n f protein LG

Several colonies from a culture plate with E. c li pHDLG were used to inoculate a preculture (LB-medium with an addition of 100 mg/l ampicillin) were cultivated at 28°C overnight. In the morning, the preculture was transferred to a larger volume (100 times the volume of the preculture) of fresh LB-medium containing ampicillin (100 mg/l) and was cultivated in vibrating flasks (200 rpm), (or fermenters) at 28°C. When an absorbence value of 0.5 was reached at 620 nm, the cultivation temperature was raised to 40°C (induction of transcription). The cultivation process was then continued for 4 hours (applies only to cultivation in vibrated flasks). The bacteria were centrifuged down upon termination of the cultivation process. The bacteria were then lysed at 4°C in accordance with an osmotic shock method (Dalböge et al., 1989). The lysate was adjusted to a pH of 7. Remaining bacteria rests were centrifuged down and the supernatent then purified on IgG-sepharose, in accordance with the protocol earlier described with reference to protein G and protein L. (Sjöbring et al., 1991, Kastern et al., 1992).

The expression system gave about 30 mg/l of protein LG when cultivation in vibrated flasks. A deposition has been made at DSSM, Identification Reference DSSM <u>E. coli</u> LE392/pHDLG.

30 Example 3

Analysis of the binding properties of protein LG

Western Blot

Protein G (the C1DC2-fragment), protein L (f ur B-

1 mm

5

10

15

20

25

domains) and protein LG were isolated with SDS-PAGE (10% acrylamide c ncentration). The isolated proteins were transfered to nitrocellulose membranes in three similar copies (triplicate). Each of these membranes was incubated with radioactively marked proteins (20 ng/ml: one of the membrane-copies was incubated with human polyclonal IgG, another with human IgG Fc-fragment and the third with isolated human IgG χ chains. Non-bound radioactively marked proteins were rinsed off and all filters were then exposed to X-ray film.

Slot-blot

Human polyclonal Ig-preparations and Ig-fragments were applied with the aid of a slot-blot appliances on nitrocellulose filters in given quantities (see Figure 10) on three similar copies. Each of these membranes was incubated with radioactively marked proteins (20 ng/ml). One of the membrane copies was incubated with protein LG, another with protein L and the third with protein G. Non-bound radioactively marked proteins were rinsed-off and all filters were then exposed to X-ray film.

The results are shown in Figures 9 and 10.

Other binding experiments have been carried out, with the following results:

TABLE

Binding of the proteins G, L and LG to immunoglobulins.

		_ • <u></u>				
Binding protein:	G	Ka	L	К _а	LG	Ka
Immunoglobulin	-	*		·		
The second secon	on to the man	g w y bla keketak ng dan man galan alama dan d		The statement of which the	and the same of	weekle e see
Human:		65 (40)		9.0	+	20
Polyclonal IgG	+	67 (10)	+	9.0		20
IgG subclasses	+	2.0	+		+	
IgG IgG ₂	+	3.1	+ .		+	
19G-2			+		+	
IgG ₃	+	6.1			+	
IgG ₄	+	4.7	+			
IgG fragment *	-					
Fc		+ 6.0 (0				т.
F(ab') ₂ *	+	0.4 (0.2)	+ (+	
kappa			+ #	1.5	+	. *
lambda	-		(-)"	•	2	
Other Ig-classes						
IgM	-		+	11.6	+	
IgA			+	10.4	+	
IgE	_		+ 1		. +	
- IgD	-	• *	· ·	·	•	
		e ⁿ in the contract of the co				
Other Species:			•			
Polyclonal		•				
Monkey +		+		+	_1	
Rabbit IgG	+	70	+	0.074	. +	
IgG-Fc	+	3.0	_		+	
IgG-F(ab')	+	0.44		• •	+	
Mouse	+	41	+	2.6	+	
Rat	+	1.5	+	0.39	+	
Goat	+	14	_		+	
GUAL	•	- ,				

TABLE (cont'd.)
Binding of the proteins G, L and LG to immunoglobulins.

Binding protein:	Ģ	Ka		L	К _а	LG	K _a
Immunoglobulin				·		. •	
			у (
Bovine IgG ₁	+	3		-		. +	
IgG ₂	+	2		-		+	
Horse	+	*		-		+	
Guinea Pig	+			+		+	
Sheep	+			-		+	
Dog	+	•				+	
Pig	+			+		+	
Hamster	+						
Cat	-			-			
Hen	-						
Monclonals &				•. ,			
Mouse	5			-		• ,	
IgG ₁	+			+	1	. +	
IgG	. +			+		+	
IgG _{2b}	+					. +	
IgG ₃	+					+ "	
IgM				+	*	+.	
IgA	-,			· +		+	
Rat							
IgG 2a	+.			+		+	
IgG	+					+	
IgG _{2C}	+			•		+	
			J				1

 K_a = affinity constant (M^{-1}). *The numerals within parenthesis disclose the affinity of a recombinant protein G comprised of two IgG-binding domains. *#A weak bond to lambda chains exists. *Binding to Pl and PLG depends on the type of light chain of Ig.

It will thus be seen that the synthesized hybrid protein LG has a broad binding activity/specificity.

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT:
 - (A) NAME: HighTech Receptro AB
 - (B) STREET: c/o Active, Skeppsbron 2
 - (C) CITY: MALMO
 - (E) COUNTRY: SWEDEN
 - (F) POSTAL CODE (ZIP): 211 20
 - (G) TELEPHONE: 040/35 07 00
 - (H) TELEFAX: 040/ 23 74 05
 - (I) TELEX: 32637 Active S
 - (ii) TITLE OF INVENTION: Hybridprotein
 - (iii) NUMBER OF SEQUENCES: 1
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25

(EPO)

- (v) CURRENT APPLICATION DATA:
 - APPLICATION NUMBER: SE PCT/SE93/00375
- (vi) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: SE 9201331-7
 - (B) FILING DATE: 28-APR-1992
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 305 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: protein
 - (iii) HYPOTHETICAL: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Escherichia coli LE392/pHDL, DSM 7054
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
- Ala Val Glu Asn Lys Glu Glu Thr Pro Glu Thr Pro Glu Thr Asp Ser 1 5 10 15
- Glu Glu Glu Val Thr Ile Lys Ala Asn Leu Ile Phe Ala Asn Gly Ser 20 25 30
- Thr Gln Thr Ala Glu Phe Lys Gly Thr Phe Glu Lys Ala Thr Ser Glu 35 40 45

Ala Tyr Ala Tyr Ala Asp Thr Leu Lys Lys Asp Asn Gly Glu Tyr Thr
50 55 60

Val Asp Val Ala Asp Lys Gly Tyr Thr Leu Asn Ile Lys Phe Ala Gly 65 70 75 80

Lys Glu Lys Thr Pro Glu Glu Pro Lys Glu Glu Val Thr Ile Lys Ala 85 90 95

Asn Leu Ile Tyr Ala Asp Gly Lys Thr Gln Thr Ala Glu Phe Lys Gly 100 105 110

Thr Phe Glu Glu Ala Thr Ala Glu Ala Tyr Arg Tyr Ala Asp Ala Leu 115 120 125

Lys Lys Asp Asn Gly Glu Tyr Thr Val Asp Val Ala Asp Lys Gly Tyr 130 135 140

Thr Leu Asn Ile Lys Phe Ala Gly Lys Glu Lys Thr Pro Glu Glu Pro 145 150 155 160

Lys Glu Glu Val Thr Ile Lys Ala Asn Leu Ile Tyr Ala Asp Gly Lys 165 170 175

Thr Gln Thr Ala Glu Phe Lys Gly Thr Phe Glu Glu Ala Thr Ala Glu 180 185 190

Ala Tyr Arg Tyr Ala Asp Leu Leu Ala Lys Glu Asn Gly Lys Tyr Thr 195 200 205

Val Asp Val Ala Asp Lys Gly Tyr Thr Leu Asn Ile Lys Phe Ala Gly
210 215 220

Lys Glu Lys Thr Pro Glu Glu Pro Lys Glu Glu Val Thr Ile Lys Ala 225 230 235 240

Asn Leu Ile Tyr Ala Asp Gly Lys Thr Gln Thr Ala Glu Phe Lys Gly
245 250 255

Thr Phe Ala Glu Ala Thr Ala Glu Ala Tyr Arg Tyr Ala Asp Leu Leu 260 265 270

Ala Lys Glu Asn Gly Lys Tyr Thr Ala Asp Leu Glu Asp Gly Gly Tyr 275 280 285

Thr Ile Asn Ile Arg Phe Ala Gly Lys Lys Val Asp Glu Lys Pro Glu 290 295 300

Glu 305

(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 921 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: unknown

- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Escherichia coli LE392/pHDL, DSM 7054
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

GCG	GTAGAAA	ATAAAGAAGA	AACACCAGAA	ACACCAGAAA	CTGATTCAGA	50
AGA	AGAAGTA.	ACAATCAAAG	CTAACCTAAT	CTTTGCAAAT	GGAAGCACAC	100
AAA	CTGCAGA	ATTCAAAGGA	ACATTTGAAA	AAGCAACATC	AGAAGCTTAT	150
GCG!	PATGCAG	ATACTTTGAA	GAAAGACAAT	GGAGAATATA	CTGTAGATGT	200
TGC	AGATAAA	GGTTATACTT	TAAATATTAA	ATTTGCTGGA	ÀAAGAAAAA	250
CAC	CAGAAGA	ACCAAAAGAA	GAAGTTACTA	TTAAAGCAAA	CTTAATCTAT	300
GCA	GATGGAA	AAACACAAAC	AGCAGAATTC	AAAGGAACAT	TTGAAGAAGC	350
AAC	AGCAGAA	GCATACAGAT	ATGCAGATGC	ATTAAAGAAG	GACAATGGAG	400
AAT	ATACAGT	AGACGTTGCA	GATAAAGGTT	ATACTTTAAA	TATTAAATTT	450
GCT	GGAAAAG	AAAAAACACC	AGAAGAACCA	AAAGAAGAAG	TTACTATTAA	500
AGC	AAACTTA	ATCTATGCAG	ATGGAAAAAC	ACAAACAGCA	GAATTCAAAG	550
GAA	CATTTGA	AGAAGCAACA	GCAGAAGCAT	ACAGATATGC	TGACTTATTA	600
GCA	AAAGAAA	ATGGTAAATA	TACAGȚAGAC	GTTGCAGATA	AAGGTTATAC	650
TTT	AAATATT	AAATTTGCTG	GAAAAGAAAA	AACACCAGAA	GAACCAAAAG	700
AAG	AAGTTAC	TATTAAAGCA	AACTTAATCT	ATGCAGATGG	AAAAACTCAA	750
ACA	GCAGAGT	TCAAAGGAAC	ATTTGCAGAA	GCAACAGCAG	AAGCATACAG	800
ATA	CGCTGAC	TTATTAGCAA	AAGAAAATGG	TAÀATATACA	GCAGACTTAG	850
AAG	ATGGTGG	ATACACTATT	AATATTAGAT	TTGCAGGTAA	GAAAGTTGAC	900
GAA	AAACCAG	AAGAATAATA	A			921

(2) INFORMATION FOR SEQ ID NO: 3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 434 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unkn wn
 - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein

- (iii) HYPOTHETICAL: NO
 - (vi) ORIGINAL SOURCE:
 (A) ORGANISM: Escherichia coli LE392/pHDLG, DSM 7055
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Ala Val Glu Asn Lys Glu Glu Thr Pro Glu Thr Pro Glu Thr Asp Ser 1 5 10 15

Glu Glu Glu Val Thr Ile Lys Ala Asn Leu Ile Phe Ala Asn Gly Ser 20 25 30

Thr Gln Thr Ala Glu Phe Lys Gly Thr Phe Glu Lys Ala Thr Ser Glu 35 40 45

Ala Tyr Ala Tyr Ala Asp Thr Leu Lys Lys Asp Asn Gly Glu Tyr Thr 50 55 60

Val Asp Val Ala Asp Lys Gly Tyr Thr Leu Asn Ile Lys Phe Ala Gly 65 70 75 80

Lys Glu Lys Thr Pro Glu Glu Pro Lys Glu Glu Val Thr Ile Lys Ala 85 90 95

Asn Leu Ile Tyr Ala Asp Gly Lys Thr Gln Thr Ala Glu Phe Lys Gly 100 105 110

Thr Phe Glu Glu Ala Thr Ala Glu Ala Tyr Arg Tyr Ala Asp Ala Leu 115 120 125

Lys Lys Asp Asn Gly Glu Tyr Thr Val Asp Val Ala Asp Lys Gly Tyr 130 135 140

Thr Leu Asn Ile Lys Phe Ala Gly Lys Glu Lys Thr Pro Glu Glu Pro 145 150 155 160

Lys Glu Glu Val Thr Ile Lys Ala Asn Leu Ile Tyr Ala Asp Gly Lys 165 170 175

Thr Gln Thr Ala Glu Phe Lys Gly Thr Phe Glu Glu Ala Thr Ala Glu
180 185 190

Ala Tyr Arg Tyr Ala Asp Leu Leu Ala Lys Glu Asn Gly Lys Tyr Thr 195 200 205

Val Asp Val Ala Asp Lys Gly Tyr Thr Leu Asn Ile Lys Phe Ala Gly 210 215 220

Lys Glu Lys Thr Pro Glu Glu Pro Lys Glu Glu Val Thr Ile Lys Ala 225 230 235 240

Asn Leu Ile Tyr Ala Asp Gly Lys Thr Gln Thr Ala Glu Phe Lys Gly 245 250 255

Thr Phe Ala Glu Ala Thr Ala Glu Ala Tyr Arg Tyr Ala Asp Leu Leu 260 265 270

Ala	Lys	Asn	Gly	Lys	Tyr	Thr 280	Ala	Asp	Leu	Glu	Asp 285	Gly	Gly	Tyr
Thr	Ile 290		Arg		Ala 295	Gly	Lys	Lys	Val	Asp 300	Glu	Lys	Pr	Glu

Glu Pro Met Asp Thr Tyr Lys Leu Ile Leu Asn Gly Lys Thr Leu Lys 305 310 315 320

Gly Glu Thr Thr Glu Ala Val Asp Ala Ala Thr Ala Glu Lys Val

Phe Lys Gln Tyr Ala Asn Asp Asn Gly Val Asp Gly Glu Trp Thr Tyr 340 345 350

Asp Asp Ala Thr Lys Thr Phe Thr Val Thr Glu Lys Pro Glu Val Ile 355 360 365

Asp Ala Ser Glu Leu Thr Pro Ala Val Thr Thr Tyr Lys Leu Val Ile 370 380

Asn Gly Lys Thr Leu Lys Gly Glu Thr Thr Thr Lys Ala Val Asp Ala 385 390 395 400

Glu Thr Ala Glu Lys Ala Phe Lys Gln Tyr Ala Asn Asp Asn Gly Val 405 410 415

Asp Gly Val Trp Thr Tyr Asp Asp Ala Thr Lys Thr Phe Thr Val Thr 420 425 430

Glu Met

(2) INFORMATION FOR SEQ ID NO: 4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1308 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Escherichis coli L392/pHDLG, DSM 7055
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

A 50	CTGATTCAGA	ACACCAGAAA	AACACCAGAA	ATAAAGAAGA	GCGGTAGAAA
100	GGAAGCACAC	CTTTGCAAAT	CTAACCTAAT	ACAATCAAAG	AGAAGAAGTA
r 150	AGAAGCTTAT	AAGCAACATC	ACATTTGAAA	ATTCAAAGGA	AAACTGCAGA
r 200	CTGTAGATGT	GGAGAATATA	GAAAGACAAT	ATACTTTGAA	GCGTATGCAG

TGCAGATAAA	GGTTATACTT	TAAATATTAA	ATTTGCTGGA	AAAGAAAAA	250
CACCAGAAGA	ACCAAAAGAA	GAAGTTACTA	TTAAAGCAAA	CTTAATCTAT	. 300
GCAGATGGAA	AAACACAAAC	AGCAGAATTC	AAAGGAACAT	TTGAAGAAGC	3.50
AACAGCAGAA	GCATACAGAT	ATGCAGATGC	ATTAAAGAAG	GACAATGGAG	400
AATATACAGT	AGACGTTGCA	GATAAAGGTT	ATACTTTAAA	TATTAAATTT	450
GCTGGAAAAG	AAAAAACACC	AGAAGAACCA	AAAGAAGAAG	TTACTATTAA	500
AGCAAACTTA	ATCTATGCAG	ATGGAAAAAC	ACAAACAGCA	GAATTCAAAG	550
GAACATTTGA	AGAAGCAACA	GCAGAAGCAT	ACAGATATGC	TGACTTATTA	600
GCAAAAGAAA	ATGGTAAATA	TACAGTAGAC	GTTGCAGATA	AAGGTTATAC	650
TTTAAATATT	AAATTTGCTG	GAAAAGAAAA	AACACCAGAA	GAACCAAAAG	70 0
AAGAAGTTAC	TATTAAAGCA	AACTTAATCT	ATGCAGATGG	AAAAACTCAA	750
ACAGCAGAGT	TCAAAGGAAC	ATTTGCAGAA	GCAACAGCAG	AAGCATACAG	800
ATACGCTGAC	TTATTAGCAA	AAGAAAATGG	TAAATATACA	GCAGACTTAG	850
AAGATGGTGG	ATACACTATT	AATATTAGAT	TTGCAGGTAA	GAAAGTTGAC	900
GAAAAACCAG	AAGAACCCAT	GGACACTTAC	AAATTAATCC	TTAATGGTAA	950
AACATTGAAA	GGCGAAACAA	CTACTGAAGC	TGTTGATGCT	GCTACTGCAG	1000
AAAAAGTCTT	CAAACAATAC	GCTAACGACA	ACGGTGTTGA	CGGTGAATGG	1050
ACTTACGACG	ATGCGACTAA	GACCTTTACA	GTTACTGAAA	AACCAGAAGT	1100
GATCGATGCG	TCTGAATTAA	CACCAGCCGT	GACAACTTAC	AAACTTGTTA	1150
TTAATGGTAA	AACATTGAAA	GGCGAAACAA	CTACTAAAGC	AGTAGACGCA	1200
GAAACTGCAG	AAAAAGCCTT	CAAACAATAC	GCTAACGACA	ACGGTGTTGA	1250
TGGTGTTTGG	ACTTATGATG	ATGCGACTAA	GACCTTTACG	GTAACTGAAA	1300
TGTAATAA			*	A unit	1308

41

Claims

1. Protein L having the ability to bind t the light chains of immunoglobulins, characterized in that the protein L has the following amino acid sequence:

					:											
	Ala 1		Glu	Asn	Lys 5	Glu	Glu	Thr	Pro	Glu 10	Thr	Pro	Glu	Thr	Asp 15	Ser
	Glu	Glu	Glu	Val 20	Thr	Ile	Lys	Ala	Asn 25	Leu	Ile	Phe	Ala	As n 30	Gly	Ser
10	Thr	Gln	Thr	Ala	Glu	Phe	Lys	Gly 40	Thr	Phe	Glu	Lys	Ala 45	Thr	Ser	Glu
	Ala	Tyr 50	Ala	туг	Ala	' As p	Thr 55	Leu	Lys	Lys	Asp	Asn 60	Glý	Glu	Tyr	Thr
15	Val 65	Asp B2	Val	λla	Хsр	Lys 70	Gly	Tyr	Thr	Leu	Asn 75	Ile	Lys	Phe	Ala	Gly. 80
	Lys	Glu	Lys	Thr	Pro 85	Glu	Glu	Pro	Lys	Glu 90	Glu	Va1	Thr	Ile	Lys 95	Ala
20	Asn	Leu	Ile	Tyr 100	Ala	Asp	Gly	Lys	Thr 105	Gln	Thr	Ala	Glu	Phe 110	Lys	Gly
	Thr	Phe	Glu 115	Glu	Ala	Thr	Ala	Glu 120	Ala	Tyr	Arg	Tyr	Ala 125	Asp	Ala	<u>L</u> eu
2.5		Lys 130	Asp	Asn	Gly	Glu	Tyr 135	Thr	Val		Val	Ala 140	Asp	Lys	Gly	Tyr
25	Thr 145	Leu	Asn	Ile	Lys	Phe 150	Ala	Gly			Lys 155	Thr	Pro	Glu	Glu	Pro 160
	Lys	Glu	Glu	Val	Thr 165	Ile	Lys	Ala	Asn	Leu 170	Ile	Tyr	Ala	λsp	Gly 175	Lys
30	Thr	Gln	Thr	Ala 180	Glu	Phe	Lys	Gly	Thr 185	Phe	Glu	Glu	Ala	Thr 190	Ala	Glu
	Ala	Tyr	Arg 195	Tyr	Ala	Asp	Leu	Leu 200	Ala	Lys	Glu	Asn	Gly 205	Lys	Tyr	Thr
25	Val	Asp 210	Val	Ala	Asp	Lys	Gly 215	туr	Thr	Leu	Asn	11e 220	Lys	Phe	Ala	Gly

432

480

528

				- B4														• •	
			Lys 225	Glu				230					Glu 235						
	5						245					230	•					Gly	
	er e c. A. a		Thr	Phe		Glu 260		Thr	Ala	Glu	Ala 265	Tyr	Arg	Tyr	Ala	Asp 270	Leu	Leu	we will
1	0		Ala	Lys	Glu 27	Asn 5	Gly	Lys	Tyr	Thr 280	Ala	Asp B5	Leu	Glu	Asp 285	Gly	Gly	Tyr	
*			Chr	Ile 290	λsn	Ile	Λrg	Phe	λla 295	Gly	Lys	Lys	Val	Asp 300	Glu	Lys	Pro	Glu	
			ilu									-			-				٠.
1	.5						n, h£v	- 2 - 7 - 7	ante	mıı	ltin	les	or 1	nixt	ures	of	the		
	į	-	ana dom	var ains	B1-	B5 h	avir	ig t	he s	ame	bind	ing	pro	pert	ies.		r		
• .			2.										i z						E .
2	20	. *					ne pr nucle						Cla	im 1	and	i has	5		5-
			CITE	101	LIOW	ing i	Iucz	-	u c 5		-		• • ,			•		**	
	*	GCG	GT	A GA	a aa	T AA	A GA	A GA	A AC	A CC	A GA	A A	CA C	CA G	AA A	CT G	AT I	'CA	∸ 8
2	25												TC T						96
													AA A						144
													AC A						192 240
:	30												AT A						288
													AA G						. 33
		AAC	TT	A AT	C TA	T GC	A GA	T GC	an - Ail	بت تتخ	-n -				-				

ACA TTT GAA GAA GCA ACA GCA GAA GCA TAC AGA TAT GCA GAT GCA TTA

ACT TTA AAT ATT AAA TTT GCT GGA AAA GAA AAA ACA CCA GAA GAA CCA

AAA GAA GAA GTT ACT ATT AAA GCA AAC TTA ATC TAT GCA GAT GGA AAA

35 AAG AAG GAC AAT GGA GAA TAT ACA GTA GAC GTT GCA GAT AAA GGT TAT

15	GAA	TAAT	'AA													•	
													,			•	921
	ACT	ATT	AAT	ATT	AGA	TTT	GCA	GGT	AAG	AAA	GTT	GAC	GAA	AAA	CCA	GAA	912
	GCA	AAA	GAA	AAT	GGT	AAA	TAT	ACA	GCA	GAC	TTA	GAA	GAT	GGT	GG <u>A</u>	TAC	864
10	ACA	TTT	GCA	GAA	GCA	ACA	GCA	GAA	GCA	TAC	AGA	TAC	GCT	GAC	TTA	TTA	816
	AAC	TTA	ATC	TAT	GCA	GAT	GGA	AAA	ACT	CAA	ACA	GCA	GAG	TTC	AAA	GGA	768
	AAA	GAA	AAA	ACA	CCA	GAA	GAA	CCA	AAA	GAA	GAA	GTT	ACT	ATT	AAA	GCA	720
ب	GTA	GAC	GTT	GCÄ	GAT	AAA	GGT	TAT	ACT	TTA	AAT	ATT	AAA	TTT	GCT	GGA	672
5	GCÀ	TAC	AGA	TAT	GCT	GAC	TTA	TTA	GCA	AAA	GAA	ААТ	GGT	AAA	TAT	ACA	624
	ACA	CAA	ACA	GCA	GAA	TTC	AAA	GGA	ACA	TTT	GAA	GAA	GCA	ACA	GCA	GAA	576

- 3. A hybrid protein, c h a r a c t e r i z e d in that it includes one or more of the B1-B5-domains according to Claim 1 which bind to the light chains in immunoglobulins of all classes, and domains which bind to heavy chains in immunoglobulin G.
- 4. A hybrid protein according to Claim 3, c h a r a c t e r i z e d in that the domains which bind to
 heavy chains in immunoglobulin G are chosen from among
 the C1- and C2-domains in protein G or from among any
 other functionally similar proteins which bind to heavy
 chains in immunoglobulin G, and variants, subfragments,
 multiples or mixtures thereof having the same binding
 properties.
- 5. A hybrid protein according to Claim 4, c h a r a c t e r i z e d in that the hybrid protein has the following amino acid sequence:

																_
	Ala i	∵al	Glu	Asn	⊥∵s 5	Glu	Glu	Thr	Pro	Glu 10	Thr	Pro	Gl _u	Thr	Asp 15	Ser
5	Glu	Glu	Glu	Val 20	Thr	Ile	Lys	Ala	Asn 25	Leu	Ile	Phe	Ala	Asn 30	Gly	Ser . ,
erd je	Thr	Gln	Thr 35	Ala	Glu	Phe	Lys	Gly 40	Thr	Phe	Glu	Lys	Ala 45	Thr	Ser	Glu
10	Ala	Tyr 50	Ala	Tyr	Ala	Asp	Thr 55	Leu	Lys	Lys	Asp	Asn 60	Gly	Glu	Tyr	Thr
.:	65					70					Asn 75					
15	Lys	Glu	Lys	Thr	Pro 85	Glu	Glu	Pro	Lys	Glu 90	Glu	Val	Thr	Ile	Lys 95	Ala
	Asn	Leu	Ile	Tyr 100	Ala	Asp	Gly	Lys	Thr 105	Gln	Thr	Ala	Glu	Phe 110	Lys	Gly
. ,			115		-		•	120				1, -	129			Leu
20		130			-		135					140				Tyr
	Thr 145	Leu	Asn	Ile	Lys	Phe 150	Ala	Gly	Lys	Glu	Lys 155	Thr	Pro	Glu	Glu	Pro 160
25	Lys	Glu	Glu	Val	Thr 165	Ile	Lys	Ala	Asn	Leu 170	Ile	Tyr	Ala	Asp	Gly 175	Lys
	Thr	Gln	Thr	Ala 180	Glu	Phe	Lys	Gly	Thr 185	Phe	Glu	Glu	Ala	190	Ala	Glu
30	Ala	Tyr	Arg 195			Asp			Ala	Lys	Glu	Asn	Gly 205	Lys	Туг	Thr
	Val	Asp 210	Val	Ala	Asp	Lys	Gly 215	Tyr	Thr	Leu	Asn	11e 220	Lys	s Phe	. Ala	a Gly
	Lys 225	Glu	Lys	Thr	Pro	Glu 230	Glu	Pro	Lys	Glu	Glu 235	Val	Thi	Ile	. Lys	240
35	Asn	Leu	Ile	Tyr	Ala 245	Asp	Gly	Lys	Thr	Gln 250	Thr	Ala	Glu	ı Phe	25	s Gly

Glu Met

	Thr	Phe	Ala	Glu 260		Thr	Ala	Glu	Ala 265		Arg	Туг	Ala	Ası 27	p Lei	ı Le
5	Ala	Lys	Glu 27		Gly	Lys	Tyr	Thr 280		Asp	Leu	Glu	Asp 285	Gly	y Gl	у Ту
:	Thr	Ile 2 90	Asn	Ile	Arg	Phe	Ala 295	Gly	Lys	: Lys	: Val	300	Glu	Ly	s Pr	o Gl
10	31u 305		<u> Met</u>	Asp	Thr	Tyr 310	Lys	Leu	Ile	Leu	Asn 315	31;	Lys	Thr	Leu	Lys 320
	Gly	Glu	Thr	Thr	Thr 325	Glu	Ala	Val	Asp	Ala 330	Ala	Thr	Ala	Glu	Lys 335	Val
15	. Phe	Lys	Gln	Tyr 340	Ala	Asn	Asp	Asn	Gly 345	Val	Asp	Gly	Glu	Trp 350	Thr	Tyr
,	Asp	Asp	Ala 355	Thr	Lys	Thr	Phe	Thr 360	Val	Thr	Glu	Lys	Pro 365	Glu	Val	Ile
	Asp	Ala 370	Ser	Glu	Leu	Thr	Pro 375	Ala	Val	Thr	Thr	Tyr 380	Lys	Leu	Val	Ile
20	Asn 385	Gly	Lys	Thr	Leu	Lys 390	Gly	Glu	Thr	Thr	Thr 395	Lys	Ala	Val	Asp	Ala 400
,	Glu	Thr	Ala	Glu	Lys 405	Ala	Phe	Lys	Gln	Tyr 410	Ala	Asn	Asp	Asn	Gly 415	Val
25	Asp	Gly	Val	Trp 420	Ťhr	Tyr	Asp	Asp	Ala 425	Thr	Lys	Thr	Phe	Thr 430	Val	Thr

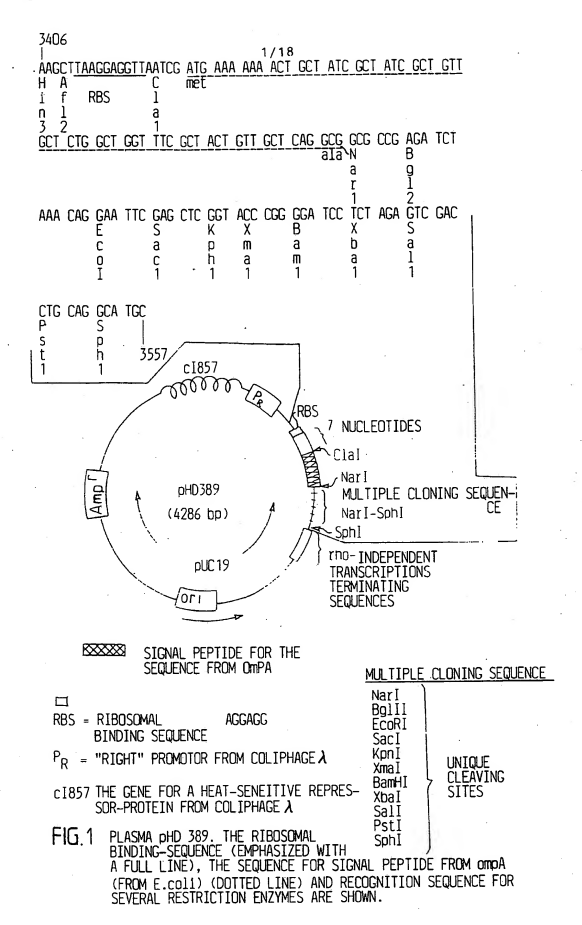
- and variants, subfragments, multiples or mixtures of the domains B1-B5 having the same binding properties.
- 6. DNA-sequence, characterized in that it codes for a protein according to Claim 5 and has the following nucleotide sequence:

					_				ac:	63.	101	CC3	CAA	ኔ.ሮጥ	GAT	TCA	∔8
						•									GAT		
٠								• • •							GGA		96
	ACA	CAA	ACT	GCA	GAA	TTC	AAA	GGA	ACA	TTT	GAA	AAA	GCA	ACA	TCA	GAA	144
	GCT	TAT	GCG	TAT	GCA	GAT	ACT	TTG	AAG	AAA	GAC	AAT	GGA	GAA	TAT	ACT	192
5	GTA	GAT	GTT	GCA	GAT	AAA	GGT	TAT	ACT	TTA	AAT	ATT	AAA	TTT	GCT	GGA	240
two to the	AAA	GAA	AAA	ACA	CCA	GAA	GAA	CCA	AAA	GAA .	GAA	GTT	ACT	ATT	AAA	GCA	288
rake etrility	AAC	TTA	ATC	TAT	GCA	GAT	GGA	AAA	ACA	CAA	ACA	GCA	GAA	TTC	AAA	GGA	336
10															GCA		384
															GGT		432
															GAA		480
															GGA	•	528
15															GCA		576
1					-										TAT		624
													:		GCT		672
							•								AAA		720
20												••					768
											•		-			GGA	816
										***					TTA		
25														-	GG <u>A</u>		864
																GAA	
					-											AAA	960
• •	GGC	GAA	ACA	ACT	ACT	GAA	GCT	GTT	GAT	GCT	GCT	ACT	GCA	GAA	AAA	GTC	1008
30	TTC	AAA	CAA	TAC	GCT	AAC	GAC	AAC	GGT	GTT	GAC	GGT	GAA	TGC	ACT	TAC	1056
	GAC	GAT	GCG	ACT	AAG	ACC	TTT	ACA	GTT	ACT	GAA	AAA	CCA	GA2	GTC	ATC	1104
	GAT	GCG	TCT	GAA	TTA	ACA	CCA	GÇC	GTG	ACA	ACI	TAC	. AAA	CT	r GTI	TTA 7	1152
																GCA	1200
35																r GTT	1248
																A ACT	
																	1308
	GAA	ATG	TAA'	LWW													

25

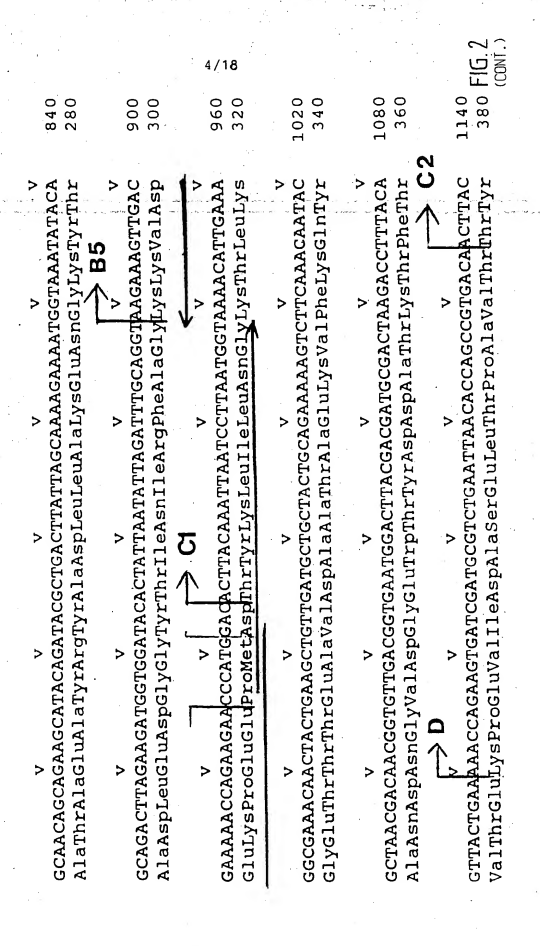
- 7. DNA-sequence, characterized in that it codes for a protein according to Claim 3 and 4.
- 8. DNA-sequence, characterized in that it hybridizes to the DNA-sequence of Claim 2, 6 or 7 under conventional conditions and codes for a protein which has the same binding properties as the protein according to any one of Claims 1 and 3-5.
- 10 9. A plasmid vector, characterized in that it includes a DNA-sequence according to any one of Claims 2 and 6-8, preferably the vector pHDLG or pHDL.
- 10. A host cell, c h a r a c t e r i z e d in that it is transformed with the hybrid plasmid according to Claim 9, in particular a host which belongs to the species <u>E. coli</u>, particularly <u>E. coli</u> LE392, or <u>Bacillus subtilis</u>. <u>Saccaromyces cerevisiae</u>, preferably Id. Ref. DSSM <u>E. coli</u> LE392 pHDL and <u>E. coli</u> LE392/pHDLG respectively.
 - 11. A method for producing a protein according to
 Claims 1 and 3-5, c h a r a c t e r i z e d by cultivating a host cell according to Claim 10 under suitable
 conditions; accumulating the protein in the culture or
 lysing the cells and extracting the protein therefrom.
- 12. A reagent apparatus for binding, separating and identifying immunoglobulins, characterized in that it includes a protein according to any one of Claims 1 and 3-5.
- 13. A composition, c h a r a c t e r i z e d in that it includes a protein according to any one of Claims 1
 35 and 3-5, and optionally additives or carriers.

14. A pharmaceutical composition, character ized in that it includes a protein according to any one of Claims 1 and 3-5, and optionally a pharmaceutically acceptable carrier or extender.



	PROTE	2/18 IN LG		*	FIG. 2 (CONT.)
20	120	180	240	300	360
GCGGTAGAAAATAAAGAAACACCAGAAACACCAGAAACTGATTCAGAAGAAGTA AlaWalGluAsnLysGluThrProGluThrProGluThrAspSerGluGluGluVal	V V V ACAATCAAAGCACACACAAACTGCAAAGGAATTCAAAGGA TTCAAAGGA TTCAAAGGA TTCAAAGGA ACAATCAAAGGA ACAATCAAAGGA ACAATCAAAGGA TTCAAAGGA TTA TTA A A CAATCAAAA TGGAAGCACACAAACTGCAGAATTCAAAGGA TTA TI I ELYSAI AAS TO TA A SA	V V V ACATTTGAAAAAGCAACATCAGAAGCTTATGCGTATGCAGATACTTTGAAGAAAGA	V V V GGAGAATATACTTTAAATTTAAATTTTGCTGGA. GGAGAATATACTGTAGATAAAGGTTATACTTTAAATTTAAATTTGCTGGA. GlyGluTyrThrValAspValAlaAspLysGlyTyrThrLeuAsnIleLysPheAlaGly	AAAGAAAAAAAGAAGCAAAAGAAGTTACTATTAAAGCAAACTTAATCTAT AAAGAAAAAAAAAA	V V V COAGAACAGCAGAATTCAAAGGAACATTTGAAGAAGCAACAGCAGAAAAAAAA

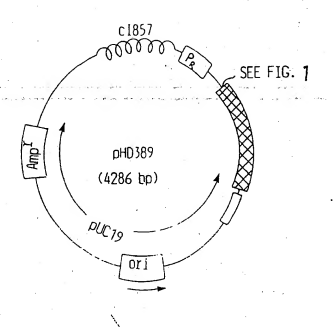
				_	- 1-
**		3/18	•	vi	F1G. 2
420 140 480 160	540 180	600	660	720	780 260 FI
CATACAGATGCAGTTAAAGAAGGACAATGGAGAATATACAGTAGACGTTGCA AlaTyrArgTyrAlaAspAlaLeuLysLysAspAsnGlyGluTyrThrValAspValAla V V V V CATAAAGGTTATACTTTAAATTTGCTGGAAAAAAAAAAA	V V AAAGAAGTTACTATTAAACTTAATCTATGCAGATGGAAAAACACAAACAGCA LysGluGluValThrIleLysAlaAsnLeuIleTyrAlaAspGlyLysThrGlnThrAla	V V V GAATTCAAAGGAACATTTGAAGAAGCAACAGAAACCATACAGATATGCTGACTTATTA CAGATATGCTGACTTATTA CAGATATGCTGACTTATTA CAGATATCAACATATATA CAGATATGCTGACTTATTA CAGATATCAAAGCAACATATA CAGATATGCTGACTTATA CAGATATCAAAAGCAACATATAA CAGATATATAA CAGATATATAA CAGATATATAA CAGATATATAA CAGATATAA CAGATATAA CAGATATAA CAGATATAA CAGATATAA CAGATATAA CATAAAA CATAAAA CATAAAAAA CATAAAAAAAA	v v v CAAAAGAAAATGGTAAATATACAGTAGACGTTGCAGATAAAGGTTATACTTTAAATATTTAAATATATAT	AAATTTGCTGGAAAAAAAAAACACCAGAAGAACCAAAAAAAA	V V AACTTAATCTATGGAAAAACTCAAACAGCAGAGTTCAAAGGAACATTTGCAGAAAAAAAA

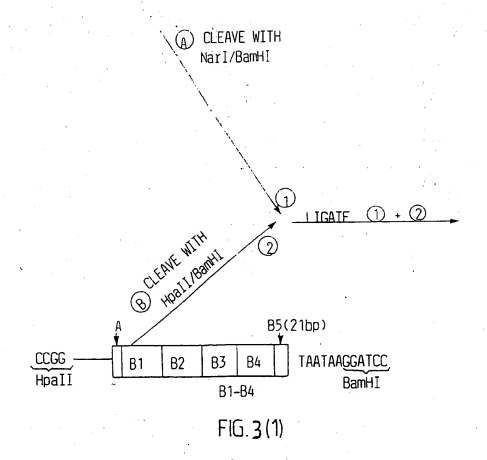


		5/18
1200	1260	F1G.2
V V AAACTTGTTATTAATGGTAAAACATTGAAAGGCGAAACAACTACTAAAAGCAGTAGACGCA LYSLeuValileAsnGlyLysThrLeuLysGlyGluThrThrThrLysAlaValAspAla	v v ordered v ordered v ordered of the control of t	V V A V A V A V A V A V A V A V A V A V

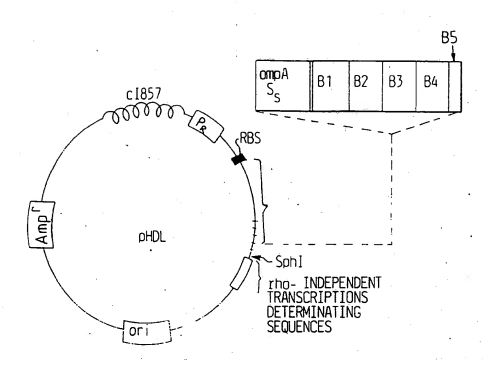
6/18

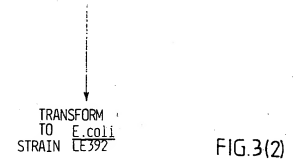
FIG.3 SCHEMATIC OVERALL VEIW OF THE PRODUCTION OF PROTEIN L

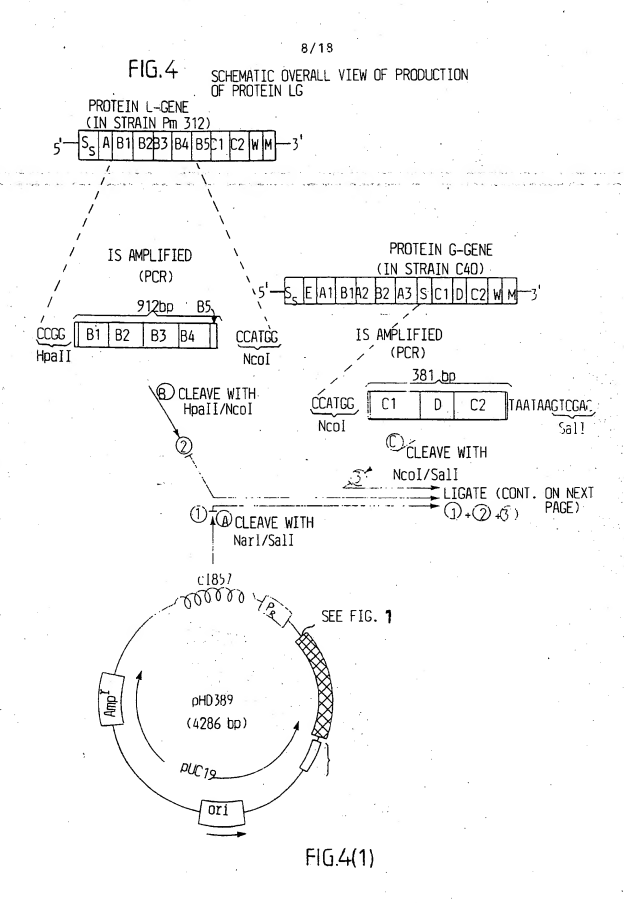




SUBSTITUTE SHEET

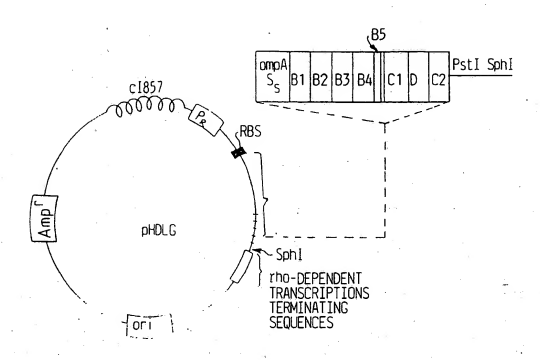


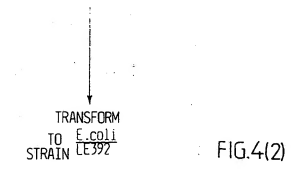


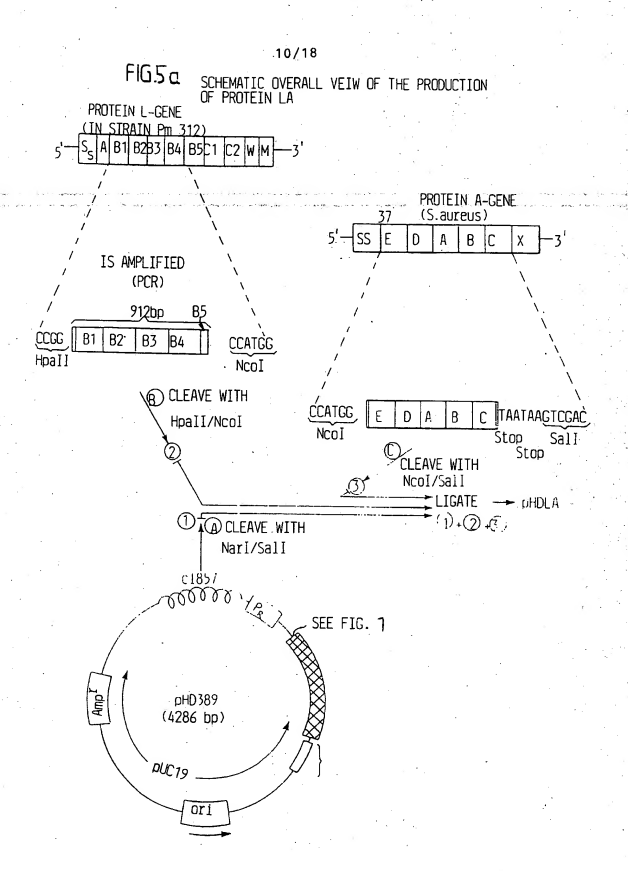


SUBSTITUTE SHEET

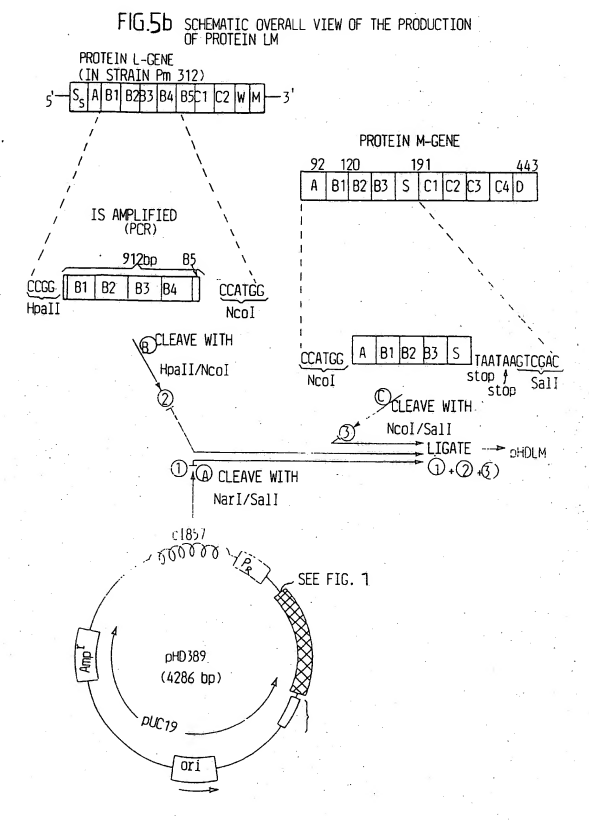
9/18

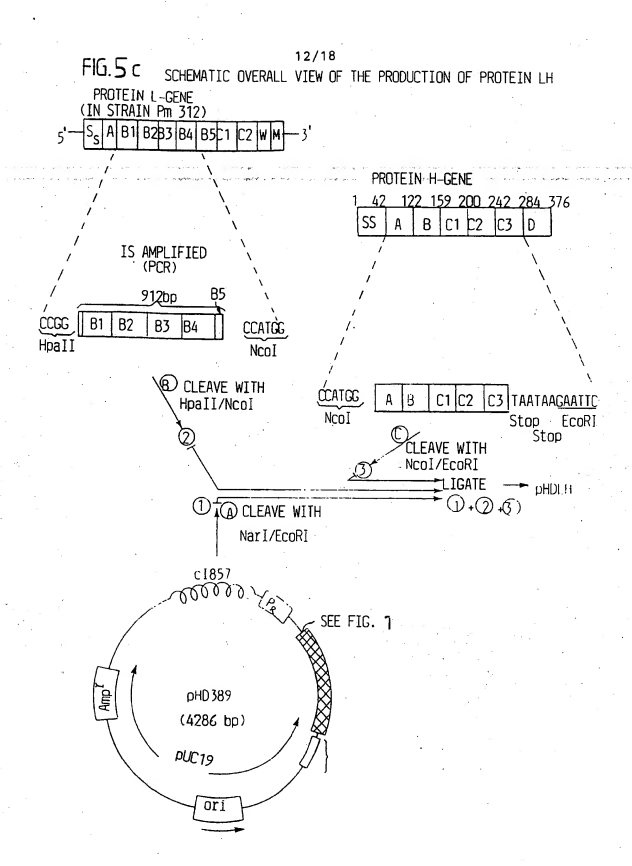


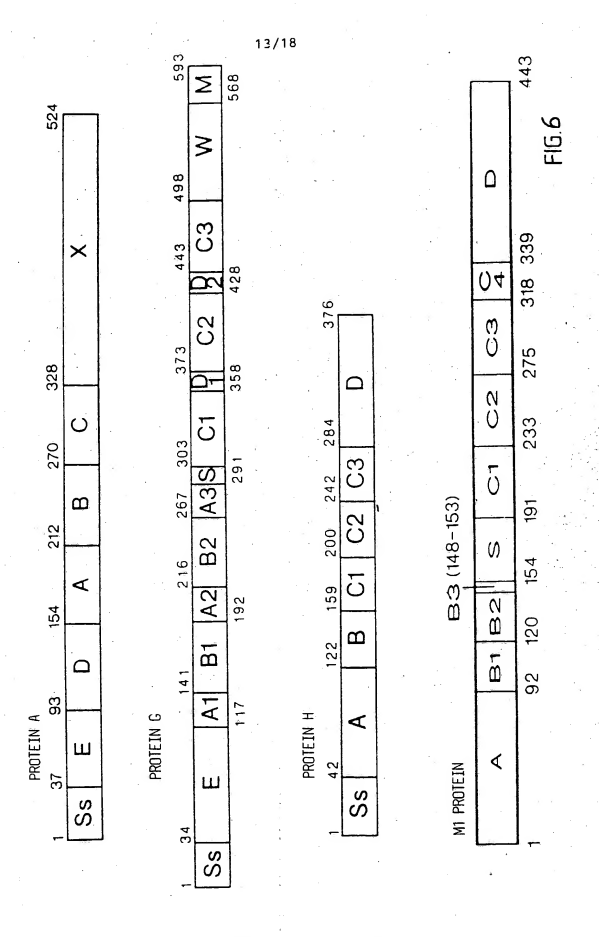




11/18







SUBSTITUTE SHEET

	(T.)	
	540	ATTAATCGTAATCTTTTAGGCAATGCAAAACTTGAACTTGATCAACTTTCATCTGAAAAA
	480	GACTATAATÄGAGCTAACGTCTTAGAAAÄGAGTTAGAAÄCGATTACTAĞAGAACAAĞAĞ Asp TyrAsnArgAl aAsnVal LeuGluLysGluLeuGluThr IleThrArgGluGluGluGlu
	420	GAAAAAGAGTTAGAAGAAAAGAAAGCTCTTGAATTAGCGATAGACCAAGCGAGTCAGGCAGTCAGGCAAAAAAGAAAAAAAA
	360	GAAGCTCTTGAATTAGCGATAGACCAGGCAAGTCGGGACTACCATAGAGCTACCGCTTTA 61 uA1 aLeuG1 uLeuA1 a I 1 eAspG1 nA1 aSer ArgAsp TyrHi sArgA1 aThrA1 aLeu
	300	CCAAAGGAATCAACAAGTTGGGATAGACAÁAGACTTGAAÁAAGAGTTAGÁAGAGAGAAAAĞ AlaLysGluSerThrSerTrpAspArgGlnArgLeuGluLysGluLeuGluGluGluLysLys
•	240 80	GAAGACCAGČGTAAAGATTYAGAAACTAAÄTTAAAAGAAČTACAACAAGAGÄCTATGACTTÄ GluAsp6lnArgLysAspLeuGluThrLysLeuLysGluLeuGlnGlnAspTyrAspLeu
	180 60	SAAGTTGCAGGAAGAGATTTTAAGAGAGCTGAAGAACTTGAAAAAGCAAAACAAGCCTTA GluValAlaGlyArgAspFheLysArgAlaGluGluLeuGluLysAlaLysGlnAlaLeu
	120	CAAAATATACGTTTACGTCACGAAAACAAGGACTTAAAAGCGAGATTAGAGAATGCAATG GlnAsnileArgLeuArgHisGluAsnLysAspLeuLysAlaArgLeuGluAsnAlaMet
	60	AACGGTGATGGTAATCCTAGGGAAGTTATÄGAAGATCTTGCAGCAAACAÄTCCCGCAATA AsnGlyAspGlyAsnFroArgGluValIleGluAspLeuAlaAlaAsnAsnFroAlaIle

3er 200	3AA 660 -75 220	3AC 720	31, 780
SAGCAGCTAACGATCGAAAAGCAAAACTTBAGGAAGAAAAAAAATCTČAGACGCAAGT 31 uG1 nLeuThr I 1 eG1 uLysA1 aLysLeuG1 uG1 uG1 uLysG1 n I 1 eSerAspA1 aSer 190	STCGTGACTTGGACGCATCACGTGAAGCTAAGAAACAGGTTGAAAAA	y y y y y y y y y y y y y y y y y y y	SCCGTCAACGGCTTCGCCGTGACTTGGACGCATCACGTGAAGCTAAGAAACAGGTT
TTBAGGAAGAAAAA	SCATCACGTGAAGC1	,	
eugluGluGluLys		ATAAGGTTAAAGAA	TGGACGCATCACGI
qo		SpLysValLysGlu	euAspAlaSerArg
TCGAAAAAGCAAAAC	GTCGTGACTTGGACG	TGACTGCTGAACTTG	GGCTTCGCCGTGACT
1 eG1 uLysA1 aLysL	rgArgAspLeuAspA	euThrAlaGluLeuA	rgLeuArgArgAspL
GAGCAGCTAACGA	CGTCAAAGCCTTCG	SATTTAGCAAACTTG	GCAAGCCGTCAAC(
GluGlnLeuThr I	ArgGlnSerLeuArg	AspLeuAlaAsnLeu	AlaSerArgGlnAr

F16.7 (CONT.)

15/18

Amino acid sequence and nucleic acid sequence for protein M1, IgG-binding somewhere between amino acid 1–190.

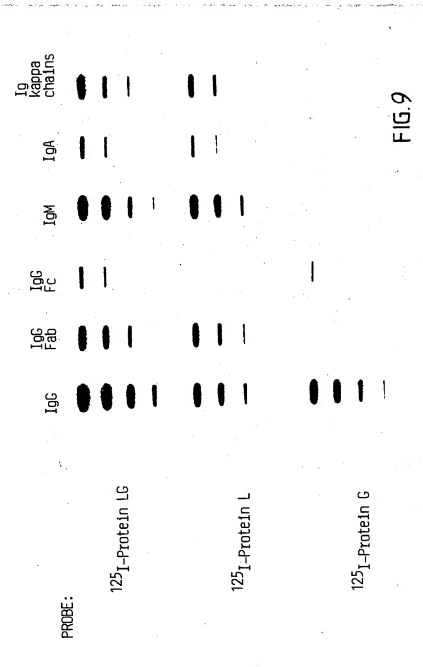
				16	5/18				
087	300	960 320	1020 340	1080 360	1140 380	1200	1260 420	1320 440	
GAAAAAGATTTAGCAAACTTGACTGAACTTGATAAGGTTAAAGAAGAAAAAAAA	, rcagacgcadaccatcaacgccttcaccttggacgcatcacgtgaagctaagaaa seraspalaseraryGlnAryLeuAryAryAspLeuAspAlaserAryGluAlaLysLys	CAAGTTGAAAAGCTTTAGAAGCAAACAACAGCAAATTAGCTGCTCTTGAAAAACTTAAC GlnValGluLysAlaLeuGluGluAlaAsnSerLysLeuAlaAlaLeuGluLysLeuAsn	AAAGAGCTTĞAAGAAAGCAAGAAATTAACAGAAAAAAGAAAAG	CTTGAAGCAGAAGCAAAGCACTCAAAGAACAATTAGCGAAACAAGCTGAAGAACTCGCA LeuGluAlaGluAlaLysAlaLeuLysGluGlnLeuAlaLysGlnAlaGluGluLeuAla	AAACTAAGAGCTGGAAAAGCATCAGAACTCACAAACCCCTGATACAAAACCAGGAAACAAAAAAAA	GCTGTTCCAGGTAAAGGTCAAGCACCAGGTACAAAACCTAACCAAAACAAAGCA AlaValFroGlyLysGlyGlnAlaProGlnALaGlyThrlysFroAsnGlnAsnLysAla	CCAATGAAGGAAACTAAGAGACAGTTACCATCAACAGGTGAAACAGCTAACCCATTCTTC PrometLysGluThrLysArgGlnLeuProSerThrGlyGluThrAlaAsnProPhePhe	V ACAGCGGCACGCGTTACTGTTATGGCAACAGCTGGAGTAGCAGTTGTAAACGCAAA ThrAlaAlaArgValThrValMetAlaThrAlaGlyValAlaAlaValValLysArgLys	GAAGAAACTAA 1329 GluGluAsn>>> 443

17/18

PROBE:

Ightappa

18/18



SUBSTITUTE SHEET

International application No. PCT/SE 93/00375

A. CLASSIFICATION OF SUBJECT MATTER

IPC5: C07K 13/00, C12N 15/31, C12N 15/62, A61K 37/02, C07K 3/18 According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC5: C07K, C12N, A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

SE,DK,FI,NO classes as above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

C. DOCU	MENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
. X	EP, A2, 0255497 (HIGHTECH RECEPTOR AB), 3 February 1988 (03.02.88)	1-2,8-14
		
Y	WO, A1, 8705631 (PHARMACIA AB), 24 Sept 1987 (24.09.87), see especially claim 9	3-14
1	· 	
Р,Х	The Journal of Biological Chemistry, Volume 267, No 18, 1992, William Kastern et al, "Structure of Peptostreptococcal Protein L and Identification of a Reeated Immunoglobulin Light Chain-binding Domain", pp. 12820-12825	1-2,8-14
		,
	*	
		<u> </u>

"I" later document published after the international filing date or priority
date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"X" document of particular relevance: the claimed invention cannot be
considered novel or cannot be considered to involve an inventive step when the document is taken alone
"Y" document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is
combined with one or more other such documents, such combination
being obvious to a person skilled in the art
"&" document member of the same patent family
Date of mailing of the international search report
2 9 -07- 1993
Authorized officer
l
Mikael G:son Bergstrand

X See patent family annex.

Form PCT/ISA/210 (second sheet) (July 1992)

Further documents are listed in the continuation of Box C.

INTERNATIONAL SEARCH REPORT

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

International application No. PCT/SE 93/00375

C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
х	INFECTION AND IMMUNITY, 58(1990-05):5 William Kastern et al: "Protein L, a Bacterial Immunoglobulin-Binding Protein and Possible Virulence Determinant", page 1217 - page 1222; see especially fig. 4 and 5	1-2,8-14		
Υ.		3-14		
and the property and pay are	الميامة بلا يريك والكور والمعاد ويري والمعادلة والأسوال وهي المعادل المنطقة المنطقة المنطقة والمناطقة والمناطقة الميامة بلا يعاد المعادلة ويري والمعادلة ويري والمعادلة والمعادلة والمعادلة والمعادلة والمعادلة والمعادلة والم	والمراجع والمراجع المراجع المحراث والمراجع والمراجع المراجع المراجع والمراجع المراجع والمراجع والمراجع والمراجع		
		•		
× X		•		
		. 8		
		* *		
f -				
-				
1				
-		}		
		*		
		·		
1				

INTERNATIONAL SEARCH REPORT

Information on patent family members

02/07/93

International application No.

PCT/SE 93/00375

Patent document cited in search rep			ent family ember(s)	Publication date	
EP-A2- 025	197 03/02/88	JP-A- US-A-	63032372 4876194	12/02/88 24/10/89	
0-A1- 870	531 24/09/87	DE-A- EP-A,B- SE-T3-	3783191 0262192 0262192	04/02/93 06/04/88	

					. 18	•
	*			·	· •	
					* .	
	•					
				*		
				. po a zado je je se	PLAN ARE	
・ ** - ** - * - * - * - * - * - * - * -	and the second section of the section of t	an official and the second		, who will shake the Market of Marke	San	3.42
•						
			* .			
		.*				
*	•					
•			· .			
	·					